Monodisperse magnetite nanoparticle tracers for in vivo magnetic particle imaging

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1. Introduction

Magnetic Particle Imaging (MPI) is a new biomedical imaging modality that produces real-time, high-resolution tomographic images of superparamagnetic iron oxide (SPIO) nanoparticle tracer distributions. In this study, we synthesized monodisperse tracers for enhanced MPI performance and investigated both, their blood clearance time using a 25 kHz magnetic particle spectrometer (MPS), and biodistribution using a combination of quantitative T2-weighted MRI and tissue histology. In vitro and in vivo MPI performance of our magnetic nanoparticle tracers (MNTs), subject to biological constraints, were compared to commercially available SPIOs (Resovist). Monodisperse MNTs showed a 2-fold greater signal per unit mass, and 20% better spatial resolution. In vitro evaluation of tracers showed that MPI performance of our MNTs is preserved in blood, serum-rich cell-culture medium and gel; thus independent of changes in hydrodynamic volume and fluid viscosity – a critical prerequisite for in vivo MPI. In a rodent model, our MNTs circulated for 15 min – 3x longer than Resovist – and supported our in vitro evaluation that MPI signal is preserved in the physiological environment. Furthermore, MRI and histology analysis showed that MNTs distribute in the reticuloendothelial system (RES) in a manner similar to clinically approved SPIO agents. MNTs demonstrating long-circulation times and optimized MPI performance show potential as angiography tracers and blood-pool agents for the emerging MPI imaging modality.

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1 Since MPI is truly a tracer-imaging technology, we refer to SPIO nanoparticles associated with MPI as magnetic nanoparticle tracers (MNTs) rather than contrast agents.

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signal, the magnetization reversal dynamics, which are governed by nanoparticle relaxation, must be tuned to the field frequency \( [7] \). Since nanoparticle size and size distribution determine the relaxation mechanism and distribution of relaxation times \([11]\), respectively, they must be tailored to the excitation frequency in order to optimize MPI performance. Furthermore, the optimized reversal dynamics, and thus MNT performance, must be preserved in biological environment for clinical relevance.

In this work we describe in detail the \textit{in vitro} and clinically relevant \textit{in vivo} MPI performance of our monodisperse MNTs. Specifically, we investigated the critical properties — blood circulation time, MPI signal per unit mass and biodistribution — that characterize MNTs suitable for \textit{in vivo} applications in MPI-based angiography, and furthermore, differentiate them from commercial SPIO tracers (Resovist\textsuperscript{TM}). To evaluate MPI performance, we used a magnetic particle spectrometer (MPS): a rapid and accurate method to assess the MPI-relevant imaging performance of MNTs, as pre-clinical and clinical scanners are still under development. In short, we demonstrate that these MNTs have appropriate physical and biological characteristics for immediate translational applications of MPI such as angiography.

To present a contextual framework for this work, in the following sections we provide a brief overview of the advantages of using MPI over traditional CT-angiography scans and summarize the physical ideas underpinning the novel method of MNT detection using MPI (comprehensive explanations can be found elsewhere \([1,12]\]).

1.1. MPI and medical imaging

SPIOs have a history of regulatory approval and clinical use; first introduced to the clinical market in 1995, SPIO nanoparticles were used for detecting liver lesions using T2-weighted magnetic resonance imaging. In 2009 for treatment of iron deficiency anemia in CKD patients. Due to their safe clinical history, SPIOs are also the materials of choice for development of MPI tracers. Unlike MRI, where the large magnetic moment of SPIO nanoparticles increases T2-relaxivity of nearby protons to enhance negative tissue contrast, MPI exploits the characteristic nonlinear magnetization of SPIO nanoparticles to construct high temporal (millisecond time-scales) and spatial resolution (sub-mm) images of nanoparticle distributions. Physical simulations and models \([7,15,16]\) suggest the possibility of sub-mm resolution and nanomolar sensitivity in MPI, while experimental results \([8,12]\) show that micromolar sensitivity and \( \sim 1 \) mm resolution are already attainable \([1]\). To approach the goal of sub-mm resolution and make MPI competitive with other tracer-based clinical imaging procedures such as x-ray CT, significant improvements in tracer design and optimization are required. Additionally, ICM agents are administered arterially using invasive/risky catheterization procedures and require high doses (3–10 ml of 320 mgI/ml ioxifan for imaging coronary arteries) to delineate narrow blood vessels from surrounding tissue. Since there is no obscuring tissue noise in MPI, the required dose of MNTs is significantly lower and MNTs with sufficient circulating times, as demonstrated here, can be safely administered intravenously.

1.2. SPIO nanoparticle detection in MPI

At room temperature and 100 s measurement time, magnetite \((Fe_3O_4)\) MNTs less than \( \sim 27 \) nm (dia.) are superparamagnetic, i.e. the net magnetic moment \((m)\) of an ensemble of particles is thermally excited and responds nonlinearly (Fig. 1a) to an external field \((H)\) \([17,18]\). In MPS, a modulating (AC) excitation field, \(H(t)\), of a fixed drive frequency (typically 25 kHz) and sufficient field strength (typically \(\mu_0H \sim 40 \) mT/p) to saturate MNTs is transmitted. The resulting time-varying magnetization response, \(m(H(t))\), induces a voltage in the receive coil, which is proportional to the time derivative of magnetization, \(m'(H(t))\) (Fig. 1b) (The constants of proportionality relating \(m'\) \([Am^2/s]\) to the measured voltage are the coil sensitivity, \(S\) \([1/m]\), and \(\mu_0\), \(4\pi \times 10^{-7}\) \([Vs/Am]\)). Fig. 1c is a parametric plot of \(m'(H(t))\) vs. \(H(t)\); since this figure is unique to each tracer formulation, we call it the Particle Response Function (PRF). For a given excitation field frequency there is a single particle diameter associated with a resonant, or perfectly in-phase, magnetization response of maximum signal intensity. Thus, it is critical that the size distribution be monodisperse for the entire ensemble of nanoparticles to magnetize simultaneously. The PRF signal intensity is also linear with MNT concentration; a critical property for \textit{in vivo} quantification of MNTs during real-time MPI scans.

In a 2-D or 3-D MPI scanner, a field gradient \((\sim 2\sim 8\) T/m) within the region of interest (ROI) creates a single point with zero-field called the field-free point (FFP) (see Refs. \([1\sim 3]\) for more details). Due to the field gradient, MNTs located in the ROI experience magnetic saturation everywhere except the FFP. Saturated MNTs do not respond to the AC-field, while MNTs within the FFP generate a time-varying signal, proportional to \(m'(H(t))\), that reveals their location. The volume of the FFP determines the instrument’s viewing window within which MNTs are resolved in space; large field gradients result in small FFP. Together, the PRF and the field gradient strength determine the system Point Spread Function (PSF); critically, the PSF is a property of the tracer and hardware. The full width half maximum (FWHM) of the PSF is the system spatial resolution and the height is the signal intensity (in V/g Fe). To generate an image, the FFP is shifted in 3-dimensions using electromagnets and a spatial distribution of MNTs in the ROI is revealed. In the \(x\)-space approach \([19]\), image reconstruction is achieved by a convolution of the tracer distribution and PSF.

![Fig. 1. Schematic illustration of magnetic nanoparticle detection in MPS. (a) The characteristic non-linear magnetization response of superparamagnetic nanoparticles to an applied field. (b) MPS uses an AC-field, \(H(t)\), to magnetize MNTs, \(m(H(t))\), and induce a time-varying signal, \(m'(H(t))\), in the receive coil. The resulting particle response function (PRF), shown in (c), is a fundamental property of MNTs.](image-url)
Alternatively, in the fast System Matrix approach, image reconstruction is achieved through matrix inversion of system function measurements\(^2\) [20]. In principle, MPS is a 0-dimensional MPI scanner (no field gradients) that measures the instrument-independent MNT performance, or the PRF, and enables tracer development independent of scanner development. Without an imaging scanner, tracer development for clinical applications of MIP is challenging. In this study, we adopted MPS to study the in vitro MPI performance of MNTs, and further evaluate their in vivo performance and bioavailability (circulation time) in a rodent model. Since all commercial scanners under development will operate at ~25 kHz, all our measurements were done in a 25 kHz MPS to align our optimization efforts for future clinical scanners.

2. Materials and methods

2.1. Synthesis of magnetite (Fe\(_3\)O\(_4\)) MNTs

Magnetite (Fe\(_3\)O\(_4\)) nanoparticles were synthesized according to published methods [21,22]. The synthesis is based on pyrolysis of the Fe\(^{3+}\) oleate precursor with excess oleic acid surfactant in 1-octadecene. Synthesis of 17 nm nanoparticles, for example, was achieved by thermal decomposition of 0.1 mmol Fe\(^{3+}\) oleate and 1.5 mmol oleic acid surfactant at a reflux temperature of 320 °C for 24 h. After synthesis, oleic acid coated MNTs were transferred from organic to aqueous phase using a PEG-ylated amphiphilic polymer (poly[maleic anhydride-alt-1-octadecene]-poly(ethylene glycol); PEGM\(_{5000}\)). Detailed synthesis, phase transfer methods and characterization results, including in vitro cytotoxicity, are discussed in previous work [23]. Resovist\(^\text{TM}\), a commercial MRI-contrast agent, was acquired from Bayer Schering Pharma AG. In addition to physical size analysis by TEM, we used the Chantrell method [24] to determine the median magnetic core diameter (D\(_m\)) of the particles and its standard deviation (σ). The latter method assumes a lognormal distribution of sizes and fits the Langevin function to the room temperature magnetization curve as measured in a vibrating sample magnetometer (VSM, Lakeshore). Hydrodynamic size was measured, both in water and in RPMI-1640 FBS medium, using dynamic light scattering (DLS). Finally, the PRF of MNTs was characterized in our home-built 25 kHz magnetic particle spectrometer (MPS) using a field amplitude of 36 mT\(_p\) (details on the design and construction of the spectrometer can be found elsewhere [8]).

For injections in mice, excess polymer and trace organic solvents were removed using sephacryl S-200 HR gel columns (GE Healthcare). Before injecting mice, MNTs were filtered with sterile 0.2-μm syringe filters and dispersed in USP-grade 1X phosphate buffered saline (PBS) at 2 gFe/L. As-received Resovist MRI-contrast agent (≈27.9 gFe/L) was also diluted to 2 gFe/L using USP-grade 1X PBS.

2.2. In vitro and in vivo studies

Female CD-1 mice [Charles River Laboratories], 25–30 g, were used for all animal studies. Sixty-five mice were used in total, forty-five for the circulation study and twenty for biodistribution. The University of Washington’s Institutional Animal Care and Use Committee approved all animal protocols.

2.2.1. In vitro evaluation

The PRF of tracers was measured using our 25 kHz MPS in biological environments, including whole blood, RPMI-1640 FBS cell-culture medium and a 1-wt% agar gel, which physically immobilized MNTs.

2.2.2. In vivo circulation study

Mice were injected with 100 μL 2.0 gFe/L MNTs dispersed in 1X PBS through the tail-vein. From each mouse, approximately 100 μL of blood was drawn retro-orbitally at two time points, after which mice were euthanized. Similar data sets were obtained for the control and Resovist groups, which were injected with 100 μL 1X PBS and 2.0 gFe/L Resovist solution, respectively. Blood samples were characterized in our 25 kHz MPS (36 mT\(_p\)), while parallel characterization in VSM was performed to compare the static (VSM) and dynamic (MPS) measurements. For MNT quantification, a series of dilutions prepared using Inductively Coupled Plasma-Optical Emission Spectrophotometer [ICP-OES, Perkin Elmer] were used to calibrate the MPS and VSM signals as a function of concentration. Whole blood samples (100 μL), extracted at various time points after tail-vein injections, were transferred in 0.5 ml Eppendorf tubes and measured directly in the MPS; no processing steps were necessary. The same samples were then transferred to polycarbonate capsules for measurements in the VSM.

2.2.3. Biodistribution

Mice were injected with either 0.5 or 2.0 gFe/L MNTs through the tail-vein. Control mice were injected with 100 μL 1X PBS. Mice were anesthetized using 5% isoflurane and maintained at 1–2% during the imaging period in the 14-T MRI Bruker Biospin. Respiration rate was monitored and maintained between 100 and 150 breaths/min. To acquire reference scans, all mice were imaged before injection. After injection, mice were imaged at 0.5-, 1-, and 24-h time points. T2-relaxivities (R2) of liver, spleen and kidneys were analyzed to quantify nanoparticle uptake. Following the 24-h time point, mice were euthanized and their organs were harvested, fixed in 10% formalin and embedded in paraffin for histological analysis. Liver, spleen, kidneys, heart, lungs and brain were collected. Tissue sections were stained for iron using Prussian blue.

2.3. Imaging parameters

A 14-T (600 MHz) vertical bore Bruker Magnetic Resonance Spectrometer was used for imaging. To acquire a T2-map, we used Bruker’s MSME-T2 protocol: 12 echo times; TR/TE = 4000/6.28...12 × 6.28. For iron oxide quantification, the T2-relaxivity as a function of concentration (r2 mmFe\(^{-1}\) s\(^{-1}\)) was measured in 1 wt. % agar gel (Fig. 2), which is often used as a tissue equivalent phantom material for measuring contrast agent relaxivity [25]. Five concentrations were used for r2 acquisition: 9.0, 17.9, 44.8, 89.5 and 179.0 μM Fe, including a 1 wt. % agarose control. For animal imaging, 25 slices, each 1-mm thick, were imaged using the sequence described above, and analyzed using ImageJ software. T2-relaxation times were calculated using the following equation:

\[ T2 = T2_0 \exp \left( \frac{-t}{\tau} \right) \]

where, T2(τ) is the MRI signal as a function of time, T2(0) is the initial signal and T2(τ) (units: s) is the relaxation time required to reach 37% of the original signal (T2(0)). The T2-relaxivity, R2 (units: s\(^{-1}\)), is the inverse of T2 and is used to estimate MNT concentration in liver, spleen and kidneys according to the following equation:

\[ T2 = T2_0 \exp \left( \frac{-t}{\tau} \right) \]
\[ R_{\text{MNT}} = R_0 + r_2*C \]  
(2)

where \( R_{\text{MNT}} \) is the T2-relaxivity after MNT injection, \( R_0 \) is T2-relaxivity before MNT injection, \( r_2 \) is the T2-relaxivity as a function of concentration (units: \( \text{mMFe}/C_0 \)), and \( C \) is MNT concentration (units: \( \text{mMFe} \)). Thus, from Equation (2), MNT concentration \( C \) is given by \( D \frac{R_2}{r_2} \), where \( D \frac{R_2}{r_2} = \frac{R_{\text{MNT}}}{R_0} \).

The relative tissue contrast before and after injections was also used for a qualitative analysis of MNT uptake in liver, spleen and kidneys. For each organ, gray scale intensities from five randomly selected regions of interest (ROI) were measured and averaged. After 24 h, mice were euthanized, their tissues harvested, and stained for iron using Prussian blue.

3. Results and discussion

3.1. Characterization of MNTs

Characterization data of the two tracers is summarized in Table 1. TEM analysis of UW MNTs (labeled as UW-17; Fig. 3a) yielded a median diameter (\( D_c \)) of 17 nm (\( \sigma = 0.15 \)), while Chantrell fitting of the same sample yielded core size of 17 nm (\( \sigma = 0.20 \)); approximately 10% smaller. Magnetization measurements typically return smaller diameters than the observed TEM size, presumably due to a magnetically “dead” surface layer. Resovist, on the other hand, showed a significant discrepancy (~65%) in the core size measurements. Extensive characterization of Resovist suggests that it is composed of two size fractions: 5 nm and 24 nm particles [26]. The larger size is a result of aggregated nanoparticles that are frequently observed in TEM images of Resovist [8]. The overall size distribution of Resovist is thus bimodal and considerably broader than UW-17. Hydrodynamic size measurements in water show that the median diameter of UW-17 is 86 nm, while Resovist is 72 nm (Fig. 3b).

In addition to the tailored size and uniform size distribution, UW-17 MNTs exhibit superior magnetic properties compared to Resovist (Fig. 3c); saturation magnetization (\( M_s \)) of UW-17 is nearly 88% of the bulk saturation value for magnetite (~470 \( \text{kA/m} \)) [27], whereas Resovist is significantly lower at only 64%. Particularly, the MPI performance of UW-17, as deduced from MPS measurements, is substantially better than Resovist (Fig. 3d). The PRF peak width (\( \Delta W_{\text{FWHM}} \)) of UW-17, which determines the instrument-independent imaging resolution and depends entirely on the tracer properties, is ~20% narrower and the signal per unit mass (\( V \text{mgFe}^{-1} \)), a critical property from a clinical dose perspective, is 2x higher than Resovist.

In our previous results [RM Ferguson et al., in preparation], we have shown that ~25 nm represents the optimum core diameter for a 25 kHz excitation frequency. In this study, however, we chose 17 nm MNTs, with a hydrodynamic diameter of 86 nm, to achieve a balance between MPI performance and pharmacokinetics. Due to strong inter-particle magnetic interactions and potential clustering of nanoparticles, hydrodynamic size of our 20 nm (and larger) MNTs is >100 nm, whereas optimal size range for improving circulation time is typically between 15 and 100 nm [28–30]. The lower and upper limits on hydrodynamic size are a result of physiological constraints: the lower limit set by the size of kidney fenestrae (~15 nm), and the upper limit by the size of sinusoidal

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**Table 1**

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<th>Characterization data of UW-17 and Resovist MNTs.</th>
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<td></td>
<td>( D_c ) [nm] (( \sigma ))</td>
</tr>
<tr>
<td></td>
<td>TEM</td>
</tr>
<tr>
<td>UW-17</td>
<td>17 (0.20)</td>
</tr>
<tr>
<td>Resovist</td>
<td>14 (0.47)</td>
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</tbody>
</table>

* TEM characterization data from [26], which provides extensive characterization of Resovist.

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**Fig. 3.** (a) TEM image of UW-17 and its corresponding histogram (inset) with a lognormal fit. Characterization of UW-17 (solid) and Resovist (dashed) MNTs in DI water: (b) hydrodynamic size measured using DLS; (c) static magnetization response measured in VSM; (d) intensity (top) and mass (bottom) normalized PRFs measured in our 25 kHz MPS.
capillaries in the liver and spleen (~50–180 nm) [31], which comprise the reticuloendothelial system (RES). Thus, particles smaller than 15 nm, and most molecular contrast agents such as ICM and Gadolinium-chelates are swiftly cleared through the kidneys, while particles larger than ~100 nm are rapidly sequestered in the RES. For effective MPI blood-pool imaging that uses a low tracer dose and does not require invasive catheterization, tracers that demonstrate both, optimal MPI performance and blood circulation characteristics are required. In the current study, UW-17 MNTs demonstrate significant improvements in MPI performance over Resovist while also exhibiting a comparable hydrodynamic size (Table 1). We are currently optimizing our phase transfer process to minimize nanoparticle clustering; these improvements will enable use of ~25 nm core and ~30–50 nm hydrodynamic diameter MNTs, which demonstrate both, excellent MPI performance and long circulation times.

3.2. In vitro tracer evaluation

In vitro evaluation of MPS performance proved critical to understanding MNT relaxation behavior in biological environments, and thus predicting their in vivo MPI performance. Blood and cell-culture medium are rich in serum proteins and simulate the angiography environment, while agar gel simulates nanoparticle immobilization in tissues. We observed clear differences in the PRF of UW-17 and Resovist MNTs (Fig. 4) relative to their corresponding PRFs in DI water. UW-17 MNTs showed no loss in MPS signal when dispersed in either blood or RPMI+10% FBS cell-culture medium, while in 1-wt% agar gel, the signal decreased marginally (~8%). Resovist, on the other hand, showed an 8% signal loss in blood, and significant loss in cell-culture medium (53%), and 1-wt% agar gel (74%).

The differences in magnetization reversal, and thus MPS performance, of UW-17 and Resovist are governed in general by two relaxation mechanisms: Néel and Brownian relaxation; each defined by a time constant $\tau_N$ and $\tau_B$, respectively. Néel relaxation...
occurs by rotation of the magnetization vector inside the particle; thus it is independent of changes in the surrounding environment. The time constant $\tau_N$ is expressed by the Arrhenius equation:

$$\tau_N = \tau_0 \exp \left( \frac{\Delta F}{k_B T} \right)$$

where $\tau_0$ is the attempt time typically in the range of $10^{-11}$–$10^{-9}$ s [32]; $k_B [1.38 \times 10^{-23} \text{ J K}^{-1}]$ is Boltzmann’s constant; $T [\text{K}]$ is the absolute temperature and $\Delta F [\text{J}]$, expressed as $kV_c(1-h)^2$, is the energy barrier to achieve magnetization reversal – it depends on the anisotropy constant $K [\text{J m}^{-3}]$, a material property of either magnetocrystalline or shape origin; the core volume $V_C [\text{m}^3]$; and the reduced field $h = H/H_K$, where $H_K$ is the internal anisotropy field. Thus, Néel relaxation depends primarily on the MNT core properties and the field amplitude. Brownian relaxation, on the other hand, involves physical rotation of the entire particle; thus it is sensitive to changes in hydrodynamic size and fluid viscosity that add resistance to rotation. The time constant $\tau_B$ is given by:

$$\tau_B = \frac{3\eta V_H}{k_B T}$$

where $\eta [\text{Pa s}]$ is the carrier fluid’s viscosity and $V_H [\text{m}^3]$ is the hydrodynamic volume. During circulation, nanoparticles often undergo opsonization – protein absorption to surface – that slows Brownian relaxation due to increase in $V_H$. Furthermore, uptake in tissues can immobilize nanoparticles and physically impede Brownian relaxation. Thus, to prevent signal loss, Brownian contribution to magnetization reversal must be minimized. Our results indicate that magnetization reversal in UW-17 MNTs is independent of changes in surrounding media, and thus predominantly Néel, whereas Resovist has a significant contribution from Brownian relaxation.

3.3. In vivo circulation study

To test the in vivo MPI performance of tracers, we injected CD-1 mice with 100 μL-2.0 gFe/L of either UW-17 or Resovist via the tail-vein, and measured the MPS signal from blood samples collected at various time intervals. Calibration curves (Fig. 5) showed that the signal was linear with concentration in both MPS and VSM measurements. Thus, they were used to quantify the raw MNT signal (Fig. 6), which was normalized to mouse body weight (in kg) and plotted as a function of time (Fig. 7). Our results highlight key features that distinguish the in vivo performance of MPI-tailored MNTs (UW-17) from traditional SPIO contrast agents (Resovist). First, as UW-17 MNTs are cleared from circulation, the MPS signal, defined by the PRF peak-height (Fig. 6b), decreases with time; it takes ~5 min for the signal to reach half its initial detected value, and nearly 15 min to reach the noise floor (Fig. 7a). On the other hand, MPS signal from Resovist nanoparticles is considerably low and nearly absent past 5 min of circulation (Fig. 7b). The increase in signal detected at 5 min suggests that the injected bolus takes 0–5 min for uniform distribution, and is subsequently cleared from circulation. Second, unlike Resovist, UW-17 shows notable agreement between MPS ($f = 25 \text{ kHz}$) and VSM ($f = 1.6 \times 10^{-6} \text{ kHz}$) measurements. The latter leads us to conclude that the circulating
UW-17 MNTs are superparamagnetic, as confirmed in the static VSM measurement, but more importantly are also MPI-capable. Resovist MNTs are detected in VSM, but not at the same scale in MPS, which suggests that though nanoparticles circulate, only a fraction of Resovist MNTs are MPI-capable; indeed, the inventors of MPI hypothesized that only 3% of Resovist contributes to MPI [1].

There are several factors that affect circulation time of nanoparticles, including hydrodynamic size, surface coating and charge, and opsonization dynamics [28]. While it is important that the properties affecting circulation time of MNTs are optimized, it is also critical that the physical properties optimizing MPI response are preserved. Our *in vitro* study shows that the MPI response of UW-17 is predominantly governed by Néel relaxation; thus independent of changes in the environment and preserved *in vivo*. We anticipate further improvement in circulation time will entail simultaneous optimization of the surface coating and the core. For

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**Fig. 7.** (a) UW-17 and (b) Resovist concentration in mouse blood (average of three runs) as a function of time.

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**Fig. 8.** Image analysis of mice injected with (a) 1X PBS, (b) 0.5 gFe/L and (c) 2.0 gFe/L UW-17 MNTs; relative contrast in T2-weighted images (TE = 6.28 ms) is the ratio of gray intensity value in ROI relative to the vertebrae in the same slice. (d and e) T2-plots for measuring change in relaxivity (∆R2) were used to quantitatively determine MNT uptake in liver, spleen and kidney.
instance, decreasing hydrodynamic size to 30–50 nm, while maintaining colloidal stability in biological media, will prolong circulation time and enable use of 20 nm (or larger) MNTs that show excellent MPI performance.

### 3.4. Biodistribution

MNTs designed for MPI are tailored precisely by controlling size to respond optimally to an applied frequency. As a result, we utilized the well-established [21,22] high temperature pyrolysis of Fe(III)-oleate in organic solvents approach; it offers exceptional control over nanoparticle size and size distribution. The resulting MNTs are hydrophobic, and subsequently transferred to water phase using a PEG-based amphiphilic polymer. Since there is no clinical history of iron oxide nanoparticles synthesized in organic solvents, we performed a preliminary study to determine the biodistribution of UW-17 MNTs using a rodent model.

For qualitative biodistribution analysis, we compared the relative MRI contrast from axial slices across the abdomen (Fig. 8a–c), and the rate of MRI-signal decay (T2-relaxivity) was used for quantitative analysis (Fig. 8d and e). It should be noted that the MRI signal from mice dosed at 2.0 gFe/L UW-17 was quenched (Fig. 8e), suggesting a high degree of MNT uptake in the liver. Since the quenched signal is difficult to quantify, we included an additional group of mice that was administered a 75% lower dose (0.5 gFe/L).

At 0.5 h post-injection, both groups (0.5 gFe/L: Fig. 8b, and 2.0 gFe/L: Fig. 8c) showed an increase in relative contrast in the liver and spleen, while contrast in kidneys was similar to pre-injection state. At the 24-h endpoint, we observed a substantial signal recovery to pre-injection levels in the 0.5 gFe/L group (Fig. 8b) suggesting MNT digestion through metabolic pathways in the liver and spleen, while the 2.0 gFe/L group showed only marginal recovery (Fig. 8c). Quantitative analysis of the 0.5 gFe/L-group supported the qualitative assessment (Table 2): from 0.5 to 24 h, \( \Delta R2 \) decreases.

### Table 2

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<thead>
<tr>
<th></th>
<th>( R2_0 ) [s(^{-1})]</th>
<th>( R2_{MNT} ) [s(^{-1})]</th>
<th>( \Delta R2 = R2_{MNT} - R2_0 ) [s(^{-1})]</th>
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<tr>
<td>Liver</td>
<td>65.9</td>
<td>108.2</td>
<td>42.3</td>
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<tr>
<td>Kidney</td>
<td>30.1</td>
<td>36.7</td>
<td>6.6</td>
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<td>Spleen</td>
<td>40.4</td>
<td>50.8</td>
<td>10.4</td>
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### Table 3

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<th>( R2_{MNT} ) [s(^{-1})]</th>
<th>( \Delta R2 = R2_{MNT} - R2_0 ) [s(^{-1})]</th>
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<tr>
<td>Liver</td>
<td>76.2</td>
<td>31.2</td>
<td>-2.6</td>
</tr>
<tr>
<td>Kidney</td>
<td>33.8</td>
<td>36.7</td>
<td>3.1</td>
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<tr>
<td>Spleen</td>
<td>52.4</td>
<td>67.1</td>
<td>14.7</td>
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\( a \) T2-fit not possible due to quenched signal (see plot for liver in Fig. 8e).

\( b \) \( \Delta R2 \) values negative.

![Tissue sections collected 24 h post-injection and stained with Prussian blue (20× magnification). Control group was injected with 1× PBS. Note the naturally high concentration of ferric iron in the spleen control makes qualitative comparison between control and MNT group difficult.](image-url)
58.6% (42.3 s\(^{-1}\)–17.5 s\(^{-1}\)) in the liver and 54.8% (10.4 s\(^{-1}\)–4.7 s\(^{-1}\)) in the spleen. Kidney \(\Delta R_2\), in comparison, decreases by a significant 84.8% (6.6 s\(^{-1}\)–1.0 s\(^{-1}\)) from 0.5 h to 24 h. After 0.5 h post-injection, there was a 21.9% increase (\(\Delta R_2 = 6.6 \text{ s}^{-1}\)) in kidney \(R_2\) value, which indicates MNTs are still circulating. However, the difference in \(R_2\) values (\(\Delta R_2 = 1.0 \text{ s}^{-1}\)) between pre-injection and 24 h post-injection is less than the 5% variability in signal measurement, thus negligible. Quantitative analysis of the 2.0 gfe/L group is summarized in Table 3; liver signal in the 2.0 gfe/L group was quenched and thus not quantifiable. However, signal in the spleen was quantified and showed an increase in \(\Delta R_2\) from 0.5 h to 24 h post-injection, suggesting continued nanoparticle uptake till the 24-h endpoint. Histological analysis (Fig. 9) of tissue sections showed excellent agreement with the corresponding static (VSM: 10 kHz) measurement. The latter con

4. Conclusion

MPI is a new imaging modality that is the first to truly exploit nanoscale magnetic properties of SPIONs. However, existing commercial SPION tracer performance sub-optimally and hinders its clinical translation. Our monodisperse MNTs (UW-17) – synthesized in non-aqueous solvents and subsequently transferred to aqueous non-aqueous fluids. IEEE T Magn 1978;14(5):975–99.


