Quantifying the motion of magnetic particles in excised tissue: Effect of particle properties and applied magnetic field

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This article presents a method to investigate how magnetic particle characteristics affect their motion inside tissues under the influence of an applied magnetic field. Particles are placed on top of freshly excised tissue samples, a calibrated magnetic field is applied by a magnet underneath each tissue sample, and we image and quantify particle penetration depth by quantitative metrics to assess how particle sizes, their surface coatings, and tissue resistance affect particle motion. Using this method, we tested available fluorescent particles from Chemicon of four sizes (100 nm, 300 nm, 500 nm, and 1 \mu m diameter) with four different coatings (starch, chitosan, lipid, and PEG/P) and quantified their motion through freshly excised rat liver, kidney, and brain tissues. In broad terms, we found that the applied magnetic field moved chitosan particles most effectively through all three tissue types (as compared to starch, lipid, and PEG/P coated particles). However, the relationship between particle properties and their resulting motion was found to be complex. Hence, it will likely require substantial further study to elucidate the nuances of transport mechanisms and to select and engineer optimal particle properties to enable the most effective transport through various tissue types under applied magnetic fields.

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1. Introduction

In the fields of magnetic drug targeting [1], hyperthermia [2], and magnetic resonance imaging [3,4] there is a need to be able to transport magnetic nanoparticles (MNPs) to desired tissue locations. How MNPs move in vivo depends on their properties, the properties of the surrounding biological milieu, and on the strength of the applied magnetic field gradient. In prior work, we analyzed how the transport of MNPs in blood vessels depends on particle constitution, size, the velocity and profile of blood flow in vessels, and the strength of the applied magnetic field gradient [5,6]. In this article, we begin to address the second half of the question: how MNPs move through the tissue between blood vessels. By measuring how particle motion in freshly excised tissue depends on particle size, coating, tissue type, and the applied magnetic field gradient, we hope to provide researchers with tools for better understanding that motion in order to help select MNP designs to improve therapy [1,7–13] and diagnosis [4,13–17].

Substantial evidence indicates that particle characteristics (size, surface chemistry, volume of magnetic content) influence their motion through biological media such as mucus [18], liquids and gels [19], and brain tissue [10,11]. In mucus [18], modifying particle size and coating led to 10 fold and 10,000 fold changes in diffusion respectively. In liquids and gels [19], among a set of particles (10 nm and 50 nm size, uncoated, polystyrene coated and dextran coated), only 50 nm dextran coated particles moved, and movement in glycerol was 20 times faster than in collagen gel. In brain tissue, the width of extracellular spaces (30–64 nm) precludes liposomes above 100 nm from penetrating the brain during convection enhanced delivery [20–22]. Particle steric coating (e.g. polyethylene glycol or PEG) and charge also influence binding to cells and thereby limit or improve diffusion in brain tissue [20,21,23].

To our best knowledge, as yet there have been no experimental studies that have quantified the ability of magnetic forces to transport different types of MNPs through different tissue types. Prior research has included the study of MNP motion in liquids [19,24] and gels [19] and indicated that MNP penetration depends strongly on the characteristics of the particles and the surrounding medium. Motion of magnetic beads within in vitro cultured cells
(fibroblasts and actins) has been studied for measurement of cytoplasmic viscosity and motility [25]; however, the focus was on rheology within cells rather than passage of MNPs through tissue. Motion of particles in cells was also studied in Zhang et al. [26] to create rotation and apoptosis of cells. MNP penetration in ex-vivo human skin was studied qualitatively in Baroli et al. [27] but only for passive diffusion.

Several forces are thought to influence the motion of MNPs in tissue [28]. The magnetic force ($F_{MAG}$) is the force applied by the external magnet on the particles. Inter-particle interaction forces [29] act on magnetized particles and can lead to their agglomeration [29–31]. Tissue resistance ($F_{TR}$) is likely composed of a viscous drag force ($F_{D}$) and the binding force ($F_{S}$) due to the adhesion between particle surface coatings and the tissue microenvironment [32–37]. The contribution of these component forces and the interplay between them is complex and has been a challenge to address. As far as we know, there are no accepted mathematical theories available for adequately capturing these complexities. Hence, in this article we focus on experimentally measuring particle movement in tissue samples.

We present a simple experimental technique to quantify the motion of magnetic particles through tissue. To arrive at these first results to understand and quantify particle motion in tissue under the influence of magnetic fields, we used freshly excised tissue (ex-vivo experiments). Even though we took sensible precautions to ensure that our excised tissue samples remained as close in their properties to living tissue as possible (excision time was less than two hours, tissue was preserved in a cool environment and was stored in phosphate buffer solution), we note that our collected results may still differ from in-vivo particle behavior because even with optimal procedures excised tissue is known to differ from live tissue [38–40]. In our approach, fluorescent MNPs were placed on top of freshly excised tissue samples and a magnetic field was applied by placing a magnet under the tissue. After a set time, the tissue was fixed (flash-frozen), sliced, imaged using a fluorescent microscope, and particle movement was then quantified by processing the three-dimensional volume of particles from stacked images using standardized quantitative metrics. Experiments conducted with and without a magnetic field distinguished the effect of diffusion versus magnetic drift. Our results indicate how particle properties and tissue types can affect particle motion, under what circumstances the magnetic field is most effective at moving particles in tissue, and which particle types among those tested should be selected for efficient magnetic transport.

2. Experimental methods

We developed and implemented a methodology to measure the rate of MNP movement in fresh tissue. To do so, we placed fluorescent MNPs on top of freshly excised rat tissue and applied a high magnetic field gradient by placing a permanent magnet at a precise distance underneath the MNPs below the tissue. This magnet moved the MNPs into and partway through the tissue. The tissue was then fixed and a 3-dimensional cryostat imaging system similar to Shen et al. [41] sliced the tissues and imaged the distribution of particles in the tissue samples. The acquired imaged data was pruned and the images were stacked into a 3-dimensional volume. Standardized metrics representing the degree of particle penetration into tissue samples were used to quantitatively assess the depth of particle penetration into the tissue. Then penetration depth was tabulated to compare the effects of particle size (hydrodynamic diameter), surface coating, tissue type (brain, etc.) on MNP movement.

**Fig. 1.** A schematic of the experimental procedure. (A) Excision of an organ from a rat. (B) Excised tissue. (C) Magnetic nanoparticles (MNPs) were placed on top of the tissue sample in solution (as a ferrofluid). The permanent magnet was then applied at a prescribed distance below the tissue sample to create a calibrated magnetic force on the particles (see placement calibration illustrated in Fig. 3). (D) Resulting distribution of particles in tissue sample after 45 min. (E) The tissue was fixed in OCT (optimal cutting temperature fluid) and then sliced and imaged using an automated cryostat and a fluorescence camera. The penetration depth of the ferrofluid was then measured and quantified by a standardized metric.
liver, and kidney), and magnetic field gradient on MNP motion in tissue. A flow diagram of this process is shown in Fig. 1 below.

2.1. Materials and preparation

Long Evans Rats (obtained from Charles River) were used to obtain the tissue samples. The rats were anesthetized using Isoflurane gas and then sacrificed. All surgical and experimental procedures were approved by the University of Maryland Animal Care and Use Committee and were in accordance with NIH Guidelines on the care and use of laboratory animals.

Freshly excised organs were stored in phosphate buffer (PBS) at 4°C for about 1–2 hours until the experiments. Tissue sections of 4–6 mm thickness were prepared, embedded in liquefied gelatin in a 10 mm petri dish, and then cooled at 4°C until the tissue was immobilized.

A variety of fluorescent magnetic nanoparticles (MNPs) purchased from Chemicell GmbH were used for our experiments. In this first study, we elected to use particles from Chemicell because we have used ChemiCell particles for many years, and are familiar with the handling, properties and behaviors of these particles. Four available sizes (100 nm, 300 nm, 500 nm and 1 μm) and four coatings (Chitosan, Starch, Lipid and PEG/P) were selected. Chemicell GmbH provided us with two types of particles. (A) NanoscreenMag particles with hydrodynamic size <300 nm and with the fluorescent dye outside the iron core beneath the external coating; (B) ScreenMag (or SiMag) particles with hydrodynamic size between 500 nm–1 μm where the fluorescent dye is incorporated in a silica shell around the core and the particle coating is around the silica shell. All particles contained a fluorescent red dye (lipophilic fluorescence dye Lumigen-Red for nanoscreenMAG and nil-blue for screenMAG) for easy visualization of the distribution of the particles in tissue. These two types of particles are illustrated in Fig. 2. In our experiments, the dye was excited at 578 nm and emission was measured at 613 nm to quantify the distribution of particles in the tissue samples. Chemicell was unable to synthesize starch particles >500 nm, lipid particles >300 nm and PEG/P particles >200 nm due to their higher molecular weight (these particles did not remain stable at the larger sizes). Our tissue experiments were conducted for those fluorescent particles that were available from Chemicell.

To keep the experimental setup simple and compact, we employed small NdFeB, grade N42, permanent bar magnets with Nickel–Copper–Nickel triple layer coated, size 1” x 1” x 2”, magnetized through the 2 in. length with poles on 1 x 1 in. surfaces (from Applied Magnets Inc.). These magnets had a high field strength (surface field strength of ~0.4 T and core strength of 1.5 T), which created a strong maximum magnetic field gradient (~30 T/m). The magnets were placed below the tissue blocks as shown in Fig. 1C in order to effectively pull particles towards the magnet. We found that these magnets applied a sufficient magnetic field gradient to effectively move the different types of MNPs through the tissue samples. The magnets were small enough to permit a convenient experimental setup but big enough to enable careful calibration of the magnetic forces applied to the particles (discussed next).

2.2. Magnetic field and force

The magnetic force we applied on the MNPs was calibrated by measuring the magnetic field and gradient around each permanent magnet. A single ferromagnetic particle will experience a magnetic force \( F = k M [\nabla \mathbf{H} / \nabla x_k \mathbf{H} D \mathbf{H} D \mathbf{x} k] \) where \( \mathbf{H} \) is the applied magnetic field, \( \overrightarrow{M} \) is the resulting magnetization of the particle, \( [\nabla \mathbf{H} / \nabla x_k] \) is the Jacobian spatial derivatives matrix, \( \mathbf{v} \) is the spatial gradient, and \( k \) is a constant that depends on particle properties. Here the first and second expressions are equivalent by the chain rule and it can be seen that the force on a MNP depends on both the particle magnetization \( \overrightarrow{M} \) and the applied magnetic field gradient \( \mathbf{v} \nabla \mathbf{H} \). In our experiments, the applied magnetic field was strong enough to saturate the particles hence \( ||\overrightarrow{M}|| \) achieved its maximal strength of approximately 3.2 emu/g. Each magnet was placed underneath the tissue samples at a location, which produced a roughly constant magnetic motive force on the particles as they moved through the tissue, samples (please see Fig. 3).

First, the magnetic field surrounding the bar magnets was measured using a Lakeshore 460-3 Channel Gaussmeter with a measurement range from 0.03 mT to 30 T. The device has a Hall probe (MMZ-2518-UH) encased in a protective brass sleeve attached to three orthogonal unislide components (from Velmex) forming a 3-D stage. The stepper motors controlling the stages have an internal step monitor for relaying signals via serial connection to a computer. The stepper motors have a resolution of
Approximate $k$ values, for particles of various size, listed in units of $N/(A^2/m^3)$. The coefficient $k$ relates the quantity $iMVH \parallel$ shown above in Fig. 3 with the actual magnetic force on the particle by $iF = k \cdot iMVH \parallel$.

<table>
<thead>
<tr>
<th>Particle type (size)</th>
<th>100 nm</th>
<th>300 nm</th>
<th>500 nm</th>
<th>1000 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k$</td>
<td>$0.0007 \times 10^{-24}$</td>
<td>$0.0178 \times 10^{-24}$</td>
<td>$0.082 \times 10^{-24}$</td>
<td>$0.658 \times 10^{-24}$</td>
</tr>
<tr>
<td></td>
<td>$N/(A^2/m^3)$</td>
<td>$N/(A^2/m^3)$</td>
<td>$N/(A^2/m^3)$</td>
<td>$N/(A^2/m^3)$</td>
</tr>
</tbody>
</table>
2.4. Procedure

For each experiment, a known volume of fluorescent MNPs (a 4 microliter droplet that stayed within the profile area of the tissue) was placed on top of a tissue surface. The tissue had been immobilized using gelatin in a petri-dish. The petri dish was then immediately placed on top of a permanent magnet, for 45 min. The tissue was placed directly above the magnet so that the maximum magnetic field was acting on the particles. As described earlier, the magnet size had been selected so that this motive force would not change too greatly as the particles traversed the tissue samples. After magnet application, the tissue was immediately fixed by flash-freezing in liquid nitrogen. The frozen tissue was then stored at −80 °C. The fixed tissue samples were then embedded in optimal cutting temperature (OCT) fluid from Tissue-Tek Inc., for simultaneous slicing and imaging using an automated cryostat imaging system previously described in Shen et al. [41]. The automated cryostat has a single field bedded in optimal cutting temperature (OCT) fluid from Tissue-Tek Inc., for simultaneous slicing and imaging using an automated cryostat imaging system previously described in Shen et al. [41]. The tissue-OCT sample was sometimes not exactly aligned in the desired top-to-bottom orientation due to error in placing the sample on the cryostat slicer. All tissue

2.5. Post-processing of images

In order to assess the depth of MNP penetration into tissues, we analyzed the images acquired by the automated cryostat. For each slice of tissue, the fluorescent camera captures two images: one bright field image showing tissue and background and one fluorescent image showing only particles. Each image captured by the camera is stored as a matrix of pixels in unsigned integer 16 format. In case of bright field images, a zero value of the pixel represents a black color and the maximum value of 2^16 represents a white color. Similarly, in the case of fluorescent field images, a maximum pixel value represents bright fluorescence while 0 represents the absence of fluorescence and particles. The images acquired by the fluorescent camera of the automated cryostat required substantial processing in order to reliably extract a distance metric. The processing of images involved the following steps:

A. **Data cleaning:** The automated cryostat slicing generated some random images along with useful images, due to electrical noise and sometimes due to overlapping of a previous slice with the current slice. To remove all spurious images, we stacked all image data for all slices and formed a vector whose every element was the total intensity of each slice/image. Since good images have tissue and OCT region, and since OCT is white (white represents high grayscale intensity values close to 2^16) while tissue is dark gray (black represents low grayscale intensity values close to zero), we could easily filter out bad images using a median filter. Example good and bad images are shown in Fig. 4A.

B. **Image alignment:** The tissue-OCT sample was sometimes not exactly aligned in the desired top-to-bottom orientation due to error in placing the sample on the cryostat slicer. All tissue

### Table 2

<table>
<thead>
<tr>
<th>Size (nm)</th>
<th>Chitosan</th>
<th>PEG/P</th>
<th>Lipid</th>
<th>Starch</th>
<th>Mean ± standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>11.2</td>
<td>13.3</td>
<td>11.7</td>
<td>14.3</td>
<td>12.6 ± 1.4</td>
</tr>
<tr>
<td>300</td>
<td>112</td>
<td>133</td>
<td>117</td>
<td>143</td>
<td>126 ± 14</td>
</tr>
<tr>
<td>500</td>
<td>6.2 × 10^-11</td>
<td>7.41 × 10^-11</td>
<td>6.5 × 10^-11</td>
<td>8 × 10^-11</td>
<td>7 × 10^-11 ± 8 × 10^-12</td>
</tr>
<tr>
<td>1000</td>
<td>100</td>
<td>N/A</td>
<td>80</td>
<td>N/A</td>
<td>74 ± 28</td>
</tr>
</tbody>
</table>

### Table 3

<table>
<thead>
<tr>
<th>Size (nm)</th>
<th>Mean Polydispersity</th>
<th>Mean Polydispersity</th>
<th>Mean Polydispersity</th>
<th>Mean Polydispersity</th>
</tr>
</thead>
<tbody>
<tr>
<td>100 nm</td>
<td>49.9 0.99</td>
<td>119 0.528</td>
<td>267 0</td>
<td>524 1.08</td>
</tr>
<tr>
<td>Starch</td>
<td>45 1.34</td>
<td>116 0.77</td>
<td>254 0</td>
<td></td>
</tr>
<tr>
<td>Lipid</td>
<td>78 0.06</td>
<td>257 0.67</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PEG/P</td>
<td>131 0.8</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Table 4

<table>
<thead>
<tr>
<th>Charge potential</th>
<th>Chitosan</th>
<th>PEG/P</th>
<th>Lipid</th>
<th>Starch</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zeta potential</td>
<td>+34</td>
<td>–20</td>
<td>–19</td>
<td>–8</td>
</tr>
</tbody>
</table>
A. **BAD IMAGE**
- DUE TO NOISE
- DUE TO OVERLAP

B. **GOOD IMAGE**
- FILTER 26
- 0

C. **CROP OUTERMOST DARK REGION**

D. **Image Thresholding**

E. **Total Intensity**
- Depth (mm)
- Centroidal Distance

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**Fig. 4.** Image processing steps: (A) A bad image resulting from random capture of an image due to noise or overlapping of a previous slice is removed. This was a rare event, but such images were removed to clean the database. On the right, a good image that is retained is shown. In the good image, the tissue region has pixel values close to zero (dark) and OCT region has pixel values up to 216 (close to white). (B) Tissue images were sometimes not correctly aligned. A typical misalignment was 20° and usually occurred due to tissue placement error (tissues had to be placed on the cryostat quickly in order to minimize temperature changes). All images were rotated until the top surface appeared horizontal. (C) Dark background was cropped until only white OCT background and dark tissue background was visible, this was done in order to facilitate image processing. (D) The image was thresholded and the bright OCT region was assigned a complete white intensity (216) so that only tissue region was considered for particle measurement. (E) In this coloring, fluorescent particles with high intensities can be seen (red corresponds to a pixel value of 216, blue corresponds to a pixel value of zero). The sum of all intensities across a single row represents the total number of particles at that depth. The resulting pink curve (bottom right) shows the distribution of particles with depth. The centroid of this pink curve quantifies the depth of penetration of particles into the tissue sample. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

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**C. Cropping for image processing:** We then cropped the first image of each sample until only the dark gray tissue region and some surrounding white OCT regions were visible in the image. This cropped region was used for all slices of the corresponding sample. This enabled further image processing.

**D. Image thresholding:** Based on bright field images and their pixel intensities, an image intensity histogram was plotted and a threshold of intensity was selected such that the dark and bright regions could be clearly demarcated. Then, every region other than the tissue region was masked by assigning pixel values of zero.

**E. Assigning regions of interest:** Since the fluorescent image and bright field image are spatially co-registered, we masked the same region in the fluorescent image that was outside the tissue region (as in step D above) by assigning pixel values of zero. As a result, only regions inside tissue have non-zero pixel intensities. This was done for all slices to ensure that only the tissue region was considered for each particle depth measurement.

**F. Total fluorescence intensity vector along tissue depth:** Each vertical tissue image (slice) is represented by a matrix of pixel intensities with each pixel corresponding to a 35 μm × 35 μm area. The degree of fluorescence is correlated to the amount of particles in that pixel. To reduce computational burden, the intensities of particle concentrations along each horizontal row of this matrix were summed up. These sums were collected into a single vertical vector \( \mathbf{I} \) each of whose elements is the net particle intensity at that tissue depth for this single vertical slice. Plotting this vector illustrates the profile of total fluorescence intensity with tissue depth, as shown in Fig. 4E. Each single vector thus formed represents total fluorescence intensity distributed along the depth for a single slice. Summing all such vectors (so summing over all vertical slices) yields a total fluorescence intensity distributed along depth for the whole tissue block: \( \mathbf{T} = \sum_{\text{slice}} \mathbf{I} \).

**G. Auto-fluorescence removal:** In order to account for auto-fluorescence in tissue, steps A–E were performed on control samples without particles. The average of all pixel intensities from control samples was used as the auto-fluorescence intensity in that tissue type e.g. in liver, kidney or brain. This average pixel intensity of auto-fluorescence was then subtracted from the fluorescence intensity computed in step F.
H. Depth metric: In order to quantify the distance traveled by the particles in each tissue sample, we computed the centroidal distance of the particle vertical distribution. Let \( d_i \) represent the total fluorescent intensity at a depth \( d_i \) (representing the ith element of the column vector in \( F \) above). That value was normalized with respect to the maximum intensity at depth \( d_i \) from the top of tissue, and then the centroidal distance that the particles moved was computed as

\[
d_c = \frac{\sum_{i=1}^{n} (d_i - l)}{\sum_{i=1}^{n} l}
\]

This distance metric was used for quantifying the movement of the tested particle types in the various tissues.

3. Results and discussion

To begin to understand the effect of magnetic field, particle characteristics, and tissue environment on the motion of particles, we studied three types of tissue: liver, kidney, and brain, and four types of particles: starch, chitosan, lipid and PEG/P. For each tissue–particle pair, we conducted two types of experiments: (1) we let the particles passively diffuse through the tissue and (2) we held a permanent magnet (0.4 T field strength at its surface) immediately below the tissue in order to exert a maximal magnetic force on the particles at the top of the tissue. After conducting the experiments as described in Section 2, we measured the fluorescence distribution of particles in tissue samples for each case and computed the centroidal distance \( d_c \). In Fig. 5 we show representative fluorescent images to illustrate penetration of three different particle types (100 nm chitosan, 100 nm PEG/P and 300 nm lipid) in three types of tissue slices (liver, kidney, and brain) resulting from passive transport and magnetic drift. As can be seen from Fig. 5, application of the magnet increased particle motion for 100 nm particles in liver and the brain.

The degree of particle penetration into tissue samples under an applied magnetic field was then quantified for each particle type and tissue type pair. Each experiment was repeated three times \( (N=3) \) and the averaged penetration depth and its standard deviation are shown below in Fig. 6. Since our experiments were repeated only three times, we computed coefficient of variation for each experiment. The coefficient of variation was 0.25 on average with a maximum of 0.78 and a minimum of 0.038. We have tabulated the details for each experiment in the Supplementary materials.

After comparing all available fluorescent Chemcell particle types and their movement through brain, liver, and kidney tissue types, we found that the average particle penetration depth in all three tissue types fell within a range of 1.78 mm and 5.6 mm when exposed to a ~0.4 T magnetic field for 45 min. Hence, the average velocity of particle motion in liver, kidney and brain tissue was found to lie between 0.66 and 2 \( \mu \)m/s. The particle motion was slowest for 100 nm starch particles through kidney and was fastest for 1 \( \mu \)m chitosan particles in kidney.

We compared the effect of particle coatings on the magnetic drift of particles through tissue. Among all coating types, for available fluorescent particles from Chemcell, we found that chitosan particles (with +34 mV zeta potential) moved better through the liver, kidney and brain than starch particles (with −8 mV zeta potential) for particles of all sizes (100 nm to 1 \( \mu \)m). In Fig. 6, the penetration of chitosan particles (green bars) is higher than the penetration of starch particles (blue bars), except for the 300 nm size where starch particles penetrated slightly deeper than chitosan in liver tissue. Starch particles with a 1 \( \mu \)m diameter were not available from Chemcell, as mentioned in Section 2.1, and are marked by a data-not-available symbol (crossed-out square, triangle or circle).

Comparison of particle penetration due to magnetic drift versus passive diffusion in all tissue types for all particle sizes tested showed that penetration due to magnetic drift is more effective than passive diffusion for strongly positively charged particles such as chitosan (34 mV) and strongly negatively charged particles such as PEG/P (−20 mV). The computed mean and standard deviation of centroidal distance penetration by chitosan and PEG/P particles, for magnetic drift versus diffusion, is shown by green and red prisms in Fig. 7.

Apart from the above clearly discernable trends, we observed some more nuanced effects of particle size on magnetic movement through tissue. Comparison of nanoscreenMag particles (up to 300 nm in size) showed that the smallest (100 nm) chitosan particles penetrated deeper than 300 nm size chitosan, in all tissue types. Among screenMag chitosan particles (500 nm size and
above), the 500 nm sized particles penetrated deeper in the liver, penetrated to the same depth in the kidney, and penetrated less deep in the brain, as compared to their 1 μm sized counterparts. The largest available 1 μm particles with chitosan coating penetrated the deepest through kidney and brain tissues, but this was not the case for motion in liver tissue.

From Fig. 6, we can also observe that penetration of particles due to magnetic drift increased with size for starch particles. This behavior was consistent for all starch particles (sizes 100–500 nm) in all tissue types. The blue triangular prisms for kidney tissue clearly show increasing penetration depth with starch particle size, although this increase is less pronounced when comparing 300–500 nm starch particles in liver and brain tissues.

We can also compare particle penetration due to magnetic drift for the available 100 nm and 300 nm lipid particles in liver, kidney and brain. As shown in Fig. 6, the particle penetration was higher for 300 nm lipid particles compared to 100 nm lipid particles in kidney and brain tissue, as shown by the taller yellow triangular prisms and cylinders and their shorter counterparts for 100 nm particles. However, this behavior was reversed in the case of liver, as shown by the taller 100 nm yellow square bar compared to the shorter 300 nm yellow bar in the liver section in Fig. 7. Lipid particles larger than 300 nm were unavailable (as mentioned in Section 2.1) and are marked by a cross-out diagonal within the...
square, triangle, or circle shapes.

We made every effort to ensure that our results in freshly excised tissue would match in-vivo behavior as closely as possible. The time between organ excision and experiment was less than two hours, and according to accepted tissue handling procedures the tissue was preserved in cool environment \( (4^\circ \text{C}) \), and stored in phosphate buffer solution. This procedure and time interval is within the time interval studied in transplant research and research for studying the mechanical properties of tissue \[38, 55–57\]. Further, the tissue organs we selected (liver, kidney, and brain) were significantly larger than the volume of MNPs used, and only the outer surface of the tissue was exposed to the liquefied gelatin used for immobilizing the tissue. This protected the particle penetration path from the external environment to some extent, and we believe it helped ensure that the collected data will be representative of magnetic particle motion in-vivo.

4. Conclusion

An experimental method was presented to quantitatively measure the penetration depth of magnetic nano-particles (MNPs) into tissue samples under the action of an applied magnetic field. In this method, MNPs were placed on top of freshly excised liver, kidney, and brain tissue samples and were then pulled into the samples by a magnet placed underneath the tissues. The tissue samples were sliced by an automated cryostat, and the degree of MNP penetration was quantified by a centroid distance metric. Tests were conducted on available fluorescent particles from Chemillex in four sizes (100 nm, 300 nm, 500 nm, and 1 \( \mu \text{m} \)) and with four different coatings (starch, chitosan, lipoid, and PEG/P). The average particle penetration depth in all three tissue types, after a 45 minute application of a 0.4 T \( \times 1\times1\times2 \) magnet, was between 1.78 mm and 5.6 mm, which corresponds to a transport velocity between 0.66 and 2 \( \mu \text{m/s} \). We found that chitosan particles moved most effectively through all three-tissue types (as compared to starch, lipoid, and PEG/P coated particles). However, we observed many additional dependencies on particle size, coating, and tissue type, which indicate that the motion of MNPs in tissue is complex and that additional studies will be required to elucidate transport mechanisms in tissue and to engineer MNPs for optimal transport in tissue. We also stress that our data was collected in freshly excised tissue, not in-vivo in live animals, and differences between excised and living tissue may affect MNP motion in ways that are not captured by this study.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.jmmm.2015.05.069.

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