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Non-Invasive Imaging of Neuroanatomical Structures and Neural Activation with High-Resolution MRI

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INTRODUCTION

Functional magnetic resonance imaging (fMRI) was developed 20 years ago as a new technique to non-invasively measure patterns of brain activity. fMRI has many useful features. Neural activity can be made visible in fully intact subjects tested under controlled experimental settings and exposed to selectable stimulation regimes. The technique is appropriate for systems where more invasive methods cannot be used (e.g., humans), and repeated measurements of the same subject are possible. Moreover, fMRI has been successfully applied in both clinical and basic scientific studies. Yet, for most fMRI studies, interpretations of the results are often complicated by the low accuracy of measurements that are based on hemodynamic changes, the blood-oxygen-level dependent (BOLD) response, rather than neural responses (Aguirre et al., 1998). In addition, imaging artifacts may interfere with the statistical analysis and while spatial resolution (~1 mm3) of fMRI exceeds other techniques that are commonly used to measure brain activity in humans (e.g., EEG or PET), temporal resolution is low because the BOLD response peaks several seconds after neural firing has occurred (Heeger and Ress, 2002). Thus, fMRI typically provides a measurement of delayed activity in large brain regions that are difficult to characterize by other means because further exploration of the underlying cellular mechanisms requires invasive procedures which are not applicable, especially in humans (Logothetis, 2008).

For animal MRI studies, low temporal resolution of the BOLD signal is a limiting factor, but scanners that produce higher magnetic fields (7.0–11.7 T) are used to improve on the spatial resolution. Animal magnet systems are typically equipped with horizontal bores of 20–40 cm diameter, and to achieve sufficient spatial resolution in smaller animals (e.g., mice), customized “head coils” have to be used (Benveniste and Blackband, 2002). Thus, for animals with very small bodies and nervous systems such as invertebrates, magnets would have to be high-field scanners equipped with sufficient gradient strength and a very small solenoid coil.

A different approach for micro-imaging of small animals is the use of standard narrow bore (5.2 cm ID) or wide bore (8.9 cm ID) high-field nuclear magnetic resonance (NMR) spectrometers that are traditionally used for macromolecular studies. Most research institutions are equipped with high-field NMR spectrometers and upgrading to micro-imaging is much less expensive than acquisition of a new animal magnet system. Micro-imaging permits scanning of small samples and benefits from improved image quality, increased spatial resolution, and/or reduced scan times due to the improvements in signal-to-noise. Spatial resolution can be better than 100 μm in more than one dimension, and imaging at this resolution is often referred to as MR microscopy (Benveniste and Blackband, 2002). However, as with any other magnet, the bore size of NMR spectrometers is fixed and smaller compared to animal magnet systems, thus limiting the diameter of the imaged sample to a few centimeters at most (i.e., a maximum of 1 cm for the narrow-bore magnets and 3 cm for wide-bore magnets).

Different paramagnetic contrast agents have been developed to improve tissue discrimination in MR images. Most MRI agents used in clinical studies are based on chelated gadolinium which...
remains in the extracellular space (Carr et al., 1984; Caravan, 2006). However, one agent that accumulates intracellularly is manganese, a calcium analog that can enter depolarized cells through voltage-gated calcium channels. Due to the tendency of the paramagnetic manganese ions (Mn$^{2+}$) to differentially accumulate in most soft tissues, higher overall signal intensity and strongly improved contrast between structures is achieved providing for a high degree of anatomical detail (Silva et al., 2004; Van der Linden et al., 2007). Moreover, Mn$^{2+}$ ions that enter active nerve cells are transported along the axons in an anterograde direction and also trans-synaptically, which allows for investigation of neural pathways and interneuronal connections (Pautler and Fraser, 2003; Lee and Koretsky, 2004). Importantly, due to the slow clearing rate of manganese from intracellular regions, the systemic application of the neural marker can take place outside of the magnet, i.e., in an unrestrained freely behaving animal, before it is subsequently imaged (Pautler et al., 1998; Pautler and Koretsky, 2002). While manganese-enhanced magnetic resonance imaging (MEMRI) is characterized by much higher sensitivity and increased signal-to-noise ratio (SNR) as compared to BOLD fMRI, there are certain shortcomings associated with using this technique in vertebrate species. Systemic injections into the bloodstream require opening of the blood–brain barrier due to limited Mn$^{2+}$ permeability, whereas focal injections into brain regions of interest (ROIs) require invasive procedures under anesthesia (Aoki et al., 2004). Moreover, manganese at high concentrations is known to be a neurotoxin (Crossgrove and Zheng, 2004; Canals et al., 2008).

Previous studies have demonstrated that the uptake of Mn$^{2+}$ into the crayfish nervous tissue is rapid, and Mn$^{2+}$ clears out of the nervous tissue slowly before it is completely eliminated (Herberholz et al., 2004). In our current study we have extended the use of MEMRI to image localized Mn$^{2+}$ uptake in live juvenile crayfish brains for identification of stimulus-evoked neural activity. We used live juvenile crayfish as an invertebrate model as they are well suited for MEMRI studies for several reasons: (1) Crayfish can be systemically injected with manganese due to the absence of a blood–brain barrier and survive long imaging sessions (>24 h) under conditions described in this manuscript. (2) Crayfish can be imaged using standard high-field NMR spectrometers equipped with micro-imaging probes and strong gradients after they are placed into small sample tubes where they remain motionless without anesthesia. We also present detailed anatomical images of the crayfish nervous system and discuss parameters critical for successful MEMRI experiments, ranging from hardware and software requirements, to imaging sequences, to specimen preparation. Our results underscore the advantages of MEMRI in obtaining non-invasive images at high spatial resolution with increased SNRs providing a novel approach that can be used for anatomical and functional studies in other small invertebrates.

**METHODS**

In previous studies images were acquired with adult crayfish using a 4.7-T horizontal bore (35 cm) magnet and a quadrature birdcage (3.7 cm) radiofrequency (RF) coil (Herberholz et al., 2004); however we achieved superior imaging results more recently by using two different NMR micro-imaging systems that are described below.

**MICRO-IMAGING SYSTEMS**

This 600 MHz (14.1 T) Avance NMR spectrometer (Bruker, Billerica, MA, USA) is located at Georgia State University (Figure 1A) and equipped with a MICRO 5 probe (4.8 G cm$^{-3}$ A$^{-1}$) for imaging, three gradient current amplifiers (40 A), gradient water cooling system using Paravision$^{	ext{TM}}$ for data acquisition, image reconstruction, and processing. Different sized (saddle coil) exchangeable inserts (0.5–10 mm) are available.

The 400 MHz (9.4 T) Bruker DMX spectrometer is located at the Armed Forces Institute of Pathology in Rockville, Maryland. It is equipped with a MICRO 2.5 probe (2.5 G cm$^{-3}$ A$^{-1}$) for micro-imaging, three gradient current amplifiers (60 A), gradient water cooling system and Paravision$^{	ext{TM}}$ software for acquisition, image reconstruction, and processing. Exchangeable RF inserts for the MICRO 2.5 range from 2.5 to 25 mm in diameter.

**ANIMAL PREPARATION**

Juvenile crayfish (*Procambarus clarkii*) of both sexes (size: 3.6–3.7 cm rostrum to telson) were used in our experiments (Figure 1B). They were prepared for imaging sessions by cooling on crushed ice for several minutes. A cannula for systemic injections of the contrast agent was constructed by inserting a short piece (∼0.2 cm) of fused silica (ID 0.2 mm; Polymicro Technologies) into one end of 50 cm of polyethylene tubing (ID 0.28 mm, OD 0.61 mm; Becton Dickinson Co.). Red food-coloring dye (10 μl) was injected into the tube by connecting a Hamilton Microliter syringe to the other end, and the length of tubing that corresponded to the injected volume was measured and marked on the tube. Next, the Hamilton syringe was replaced by a MicroFil non-metallic syringe needle (World Precision Instruments) and connected to a 1-ml syringe. The syringe was filled with deionized water and the cannula was flushed multiple times. Next, the cannula was filled with MnCl$_2$ (manganese [II] chloride tetra-hydrate, minimum 99%; Sigma-Aldrich) dissolved in buffered (pH 7.4) crayfish saline. Once filled, a small bubble was introduced at the end of the polyethylene tubing and a 1-ml syringe filled with food-color dye was connected to the syringe needle. In this way, an easy visual reference (the position of
the dye within the transparent tubing) was available to determine the amount of Mn2+ injected into the animal. Under a microscope, a small hole was drilled with a 27-gauge needle in the dorsal carapace of the anesthetized crayfish. The location of the hole was always above the pericardial sac to the immediate left or right of the heart. The fused silica attached to the polyethylene tube was inserted into the hole. The entry of the polyethylene tube was prevented by its larger diameter. The juncture was then sealed with a few drops of superglue gel (Figure 1C, arrowhead).

Before acquiring neuroanatomical scans, the animals were injected with different concentrations of MnCl2 while performing random behavioral activities (see Results). For measurements of stimulus-dependent neural activation, the animals were implanted with a pair of fine silver wire electrodes (A-M Systems Inc.); two different implantations were performed. For stimulation of the antenna II nerve (mechanosensory input to the brain), two small holes were punctured using metallic insect pins into the fifth basal segment of the right or left antenna (posterior to the first segment of the flagellum) to allow access to the nerve (Figure 1C, arrows). Approximately 1 mm of insulation was stripped off one end of the electrodes and the bare wires (0.2 mm diameter) were inserted through the holes in the cuticle and placed next to each other onto the nerve. They were attached to the cuticle using superglue gel and the other ends were connected to a square pulse stimulator (Model SD9; Grass Technologies). For stimulation of the ventral nerve cord and activation of the lateral giant escape circuit, two small holes were punctured into the soft membrane covering the ventral side of the nerve cord between the fourth and fifth abdominal segments. The stripped-off ends of the wires were inserted through the holes and placed next to each other onto the ventral nerve cord. The electrodes were glued in place and the other ends were connected to a pulse stimulator (Model 2100; A-M Systems Inc.).

Different stimulation protocols were applied while the animals were injected with MnCl2 before the start of the imaging session (see Results). Before placing the animals into the magnet, the cannula was cut off near the entrance point into the carapace, the implanted electrodes were carefully removed using forceps, and the animals were gently moved into either transparent plastic tubes (55 mm length, 10 mm diameter) or conventional NMR tubes (180 mm length, 10 mm diameter; Figure 1D). Most animals remained motionless inside the tubes and did not need any additional restrain. However, to prevent motion artifacts during high-resolution scans, the cephalothorax of most animals that had received antenna II nerve stimulation was embedded in 2% agarose. The agarose completely eliminated movements of the antennae and eyestalks during long scans without affecting the quality of the images. After scans were completed, the agarose was removed from the animals before they were returned to their holding tanks.

**IMAGING PROTOCOLS**

The overall signal intensity in an MR image is dependent on tissue specific parameters such as the proton density (PD), the longitudinal (T1), and the transverse (T2) relaxation times. Signal is also affected by instrument-dependent parameters such as sensitivity of the electronic circuitry of the scanner, repetition time (TR), echo time (TE) as well as other factors such as inversion time and flip angle based on the specific imaging sequence used. The contrast between different tissues in a MR image is determined by the difference in the signal intensities resulting from the tissue specific parameters. Additionally, the combination of TR and TE define the “weighting” (e.g., T1, PD, T2) of the image and different weightings generate optimal results for specific tissues. In MEMRI studies, a positive contrast agent like manganese shortens the T1 relaxation time in the absorbed tissue consequently producing a bright signal in T1-weighted images (TR << T1, TE < T2).

T1-weighted images were acquired with a standard multi-slice-multi-echo (MSME) imaging sequence with a TR ranging from 600 to 1005 ms and a TE of 8 to 10.4 ms, respectively. Multi-slice T1-weighted images were acquired with a rapid acquisition with relaxation enhancement (RARE) sequence with TR = 2 s, TE = 25 ms, and a RARE factor of 4. Three-dimensional images were acquired using a RARE imaging sequence with TR/TE = 800/8 ms and a RARE factor of 8. The field of view (FOV) in combination with matrix size determines final resolution of images and affects acquisition time, whereas a higher number of averages improves SNR but increases total acquisition time. The combination of parameters selected for each scan is specified in Section “Result.”

**POST-IMAGING DATA PROCESSING**

The Paravision software available with the Bruker imaging systems can be used for image reconstruction. Image data can also be exported to a DICOM format which can then be viewed, processed, and analyzed by the other software packages. Much of the data processing of images presented in this study was performed using ImageJ (Abramoff et al., 2004). ImageJ is a public domain image processing and analysis program written in Java. It offers over 500 different available plug-ins including many that are relevant for post-imaging data analysis, e.g., measurements of intensity thresholds and grayscale values within a pre-selected image area, and it accepts image stacks that can be visualized in 3D.

We used the ImageJ software to compare manganese uptake in the brains of five stimulated samples and one unstimulated sample by measuring grayscale values for four consecutive slices (100 μm thick) that covered the major brain structures. This ensured that we covered most of the brain regions axially and allowed us to analyze the differences between the sides of the brain objectively. ROIs were specified for each sample according to image size and kept constant within each sample. Two ROIs of identical size were selected covering either the left or right half of the brain; the dimensions of the ROIs were determined by viewing a single image that showed most of the brain areas. Individual grayscale values were measured within each ROI for T1- and/or T1-weighted brain images of both the stimulated and unstimulated animal. The mean grayscale values were averaged over all four brain slices for the respective sides (left and right) and compared statistically with a non-parametric test (Wilcoxon signed-rank test) using SPSS (Version 17.0).

**RESULTS**

**NEUROANATOMY**

Traditional neuroanatomical studies use histological and microscopic techniques that provide superb spatial resolution, but are destructive and labor and time intensive. On the other hand, MRI is non-invasive and suitable for small nervous systems when sufficient contrast between different brain parts can be detected.
Michaelis et al., 2005). In our earlier study, we found that traditional MRI is not sufficient to differentiate between brain structures in crayfish, but this could be dramatically improved by systemic injections of MnCl₂ into the hemolymph. We have also previously shown that uptake of Mn²⁺ begins several minutes after administration of the contrast agent and remains in the nervous tissue for up to 48 h before it is slowly eliminated (Herberholz et al., 2004).

In our current study, we first imaged Mn²⁺ uptake into the nervous system of a juvenile crayfish (Figure 2). The animal was slowly (over 2–3 min) injected with 45 µl of 50 mM MnCl₂ and allowed to move freely for 30 min in a small water-filled but otherwise empty tank following the injection. After this period, the cannula was removed and the animal was placed in the magnet. Figure 2A shows one single section (selected from a total of 32 coronal sections) that provides a good view of both the central brain regions and the right eyestalk (left eye is out of plane). Individual neuroanatomical structures can be seen, including different ganglia located in the right eyestalk (Figure 2B), the accessory and olfactory lobes in the central brain area (Figure 2C, arrows), and the rostral parts of the brain connectives (Figures 1A,C, arrowheads). The following imaging parameters were used for this T₁-weighted scan sequence: TR = 600 ms, TE = 10.4 ms, FOV = 1.3 cm × 1.3 cm, matrix = 256 × 256, averages = 8, slice thickness = 100 µm, (in-plane) resolution = 50 µm × 50 µm, acquisition time = 21 min.

Mn²⁺ uptake into the ventral nerve cord is shown in Figure 2D. A different animal was injected with 100 µl of 20 mM MnCl₂ and allowed to move freely and undisturbed in a small tank for 15 min before the imaging session. After this period, the cannula was removed and the animal was placed in the magnet. A total of 128 sections were taken using a 3D imaging protocol (see below for parameters). Figure 2D represents a single sagittal section from the 3D stack showing the animal inside the tube with its tail curled under the abdomen. The entire ventral nerve cord as well as individual thoracic and abdominal ganglia are visualized. The following imaging parameters were used for this T₁-weighted RARE scan sequence: TR = 800 ms, TE = 8 ms, rare = 8, FOV = 2 cm × 1 cm × 1 cm, matrix = 256 × 128 × 128, averages = 32, resolution = 78 µm × 78 µm × 78 µm, acquisition time = 4 h, 33 min.

**NEURAL ACTIVITY**

Besides its important role as a contrast agent for neuroanatomical studies, we have tested the suitability of Mn²⁺ as a marker for neural activity by imaging uptake into the brain after electrical stimulation of the right or left antenna II nerve in five different animals. All five animals received the same dose of MnCl₂, received the same type of antenna II nerve stimulation, and were scanned using the same imaging protocol. An example of the neural activity after left antenna II nerve stimulation in one representative animal is shown in Figures 3 and 4. The animal was slowly injected with 40 µl of 25 mM MnCl₂ (over 4–5 min). Stimulation with electric shocks (1 V at 4 Hz) of the antenna II nerve was started simultaneously with MnCl₂ injection and continued for 25 min after the injection was finished. Immediately after stimulation, the implanted electrodes and cannula were removed and the animal was placed into the scanner. A total of 40 axial sections through the head region were taken and two individual slices (200 µm apart) are shown. Using a T₁-weighted imaging protocol, uptake of Mn²⁺ into the brain is visible in both sections (Figures 3A,B). More Mn²⁺ uptake can be seen on the left (i.e., stimulated) side of the brain, especially in the left antenna II nerve and the left accessory lobe (note that in these axial slices, the animal’s left side corresponds to the viewer’s right side). For better identification of brain structures, a schematic of the main brain regions allows to compare the histological reconstruction to our MEMRI images. Positions of both the antenna II nerve and the left accessory lobe are indicated. The imaging parameters for this scan were as follows: TR = 800 ms, TE = 10.4 ms, FOV = 0.4 cm × 0.4 cm, matrix = 128 × 128, averages = 32, slice thickness = 100 µm, (in-plane) resolution = 31 µm × 31 µm, acquisition time = 55 min.

The same animal was subsequently imaged using a T₁-weighted RARE imaging sequence (Figures 3C,D). Using this imaging protocol, Mn²⁺ uptake into the brain was clearly visible, with more contrast obtained than in the previous protocol. However, the difference in Mn²⁺ uptake between the two sides of the brain observed in the T₁-weighted images was obscured in these T₁-weighted images. The imaging parameters were as follows: TR = 2211 ms, TE = 25 ms, RARE = 4, FOV = 0.4 cm × 0.4 cm, matrix = 128 × 128, averages = 32, slice thickness = 100 µm, (in-plane) resolution = 31 µm × 31 µm, acquisition time = 38 min.
We determined individual and mean grayscale values within pre-selected ROIs that covered the left and right brain regions of this animal and a control (unstimulated) animal (Table 1). The same analysis was used to measure grayscale values in the control animal of equal size and weight (Figure 4). The control animal was injected with the same amount of MnCl₂ and scanned at the same time after injection but did not receive any stimulation of the antenna II nerve. It was scanned using the same imaging parameters as used for the stimulated animals except averages = 48 and FOV = 0.55 cm × 0.55 cm. The difference in the individual and mean grayscale values, of the left and the right side, was highest for the T₂-weighted images obtained from the animal that received a combination of MnCl₂ injection and antenna II stimulation; the difference was smaller for T₁-weighted images obtained from the same animal. In comparison the individual and mean grayscale values were similar for both sides of the brain in the control animal for the T₁ and T₂-weighted images. Thus, the differences (stimulated animal) or the lack thereof (unstimulated control animal) between the two sides of the brain is consistent throughout the 400 μm region covered axially by these slices.

In order to confirm the suitability of MEMRI for functional imaging, we obtained T₁- and T₂-weighted scans after antenna II stimulation and manganese injection in four additional animals using the 400 MHz spectrometer. We determined individual and mean grayscale values within pre-selected ROIs that covered the left and right brain regions (Table 1). The regions of interest (ROIs) for the two sides of the brain were selected using a single T₁-weighted image from each animal and kept identical for all slices. The ROIs are shown as rectangular boxes in the respective images; the red box in the image of the stimulated animal indicates the side of the animal that received electric stimulation of the antenna II nerve. The imaging parameters for each scan are given in the results.
Table 1 | Comparison of individual and mean grayscale values for one stimulated and one unstimulated sample.

<table>
<thead>
<tr>
<th>Protocol</th>
<th>Stimulated</th>
<th>Unstimulated (control)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Imaging</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Area</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Left brain half</td>
<td>Right brain half</td>
</tr>
<tr>
<td>ROI</td>
<td>1.5 cm x 1.2 cm</td>
<td>1.5 cm x 1.2 cm</td>
</tr>
<tr>
<td>Slice 1</td>
<td>130.869</td>
<td>113.377</td>
</tr>
<tr>
<td>Slice 2</td>
<td>153.981</td>
<td>138.471</td>
</tr>
<tr>
<td>Slice 3</td>
<td>151.244</td>
<td>131.287</td>
</tr>
<tr>
<td>Slice 4</td>
<td>141.001</td>
<td>124.755</td>
</tr>
<tr>
<td>Mean</td>
<td>144</td>
<td>127</td>
</tr>
<tr>
<td></td>
<td>1.2 cm x 1.1 cm</td>
<td>1.2 cm x 1.1 cm</td>
</tr>
<tr>
<td>Gray scale values</td>
<td>72.245</td>
<td>65.396</td>
</tr>
<tr>
<td>Slice 1</td>
<td>73.846</td>
<td>67.342</td>
</tr>
<tr>
<td>Slice 2</td>
<td>72.679</td>
<td>68.205</td>
</tr>
<tr>
<td>Slice 3</td>
<td>80.489</td>
<td>82.125</td>
</tr>
<tr>
<td>Slice 4</td>
<td>75</td>
<td>71</td>
</tr>
</tbody>
</table>

Both animals were imaged using the 600 MHz spectrometer. Regions of interest (ROIs) were specified for each animal according to image size and kept constant within each sample. The mean grayscale values were averaged over four slices (400 μm) covering the major regions of the brains. The intensities were measured from 8-bit grayscale images (grayscale range 0–255).

Table 2 | Comparison of mean grayscale values of brain areas ipsilateral and contralateral to the electrically stimulated antenna II nerve for all five tested individuals.

<table>
<thead>
<tr>
<th>Crayfish</th>
<th>Stimulated side</th>
<th>Unstimulated side</th>
<th>Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>144</td>
<td>127</td>
<td>17 ± 2</td>
</tr>
<tr>
<td>2</td>
<td>107</td>
<td>88</td>
<td>19 ± 3</td>
</tr>
<tr>
<td>3</td>
<td>113</td>
<td>110</td>
<td>3 ± 4</td>
</tr>
<tr>
<td>4</td>
<td>104</td>
<td>71</td>
<td>33 ± 9</td>
</tr>
<tr>
<td>5</td>
<td>101</td>
<td>92</td>
<td>9 ± 5</td>
</tr>
</tbody>
</table>

Crayfish 1 was imaged using the 600 MHz spectrometer and crayfish 2–5 were imaged using the 400 MHz spectrometer. Differences in grayscale values are shown along with standard deviations. Within an animal, the mean grayscale values were averaged over four slices (400 μm) covering the major regions of the brains. The intensities were measured from 8-bit grayscale images (grayscale range 0–255).

ganglia as well as ganglionic nerve roots can be seen. The imaging parameters for this T1-weighted scan were as follows: TR = 1005 ms, TE = 10 ms, FOV = 2 cm x 1 cm, matrix = 256 x 128, averages = 32, slice thickness = 100 μm, (in-plane) resolution = 78 μm x 78 μm, acquisition time = 1 h, 8 min.

The same animal was subsequently imaged in the axial plane. Figure 5B shows a selected single section (from 11 axial slices) of the third abdominal ganglion. The boundaries of the abdominal ganglion and what appears to be individually labeled cells within the ganglion have been visualized by highlighting the most intense labeled areas (Figure 5C). We used the following imaging parameters for this T1-weighted scan: TR = 805 ms, TE = 10 ms, FOV = 0.5 cm x 0.5 cm, matrix = 128 x 128, averages = 64, slice thickness = 100 μm, resolution = 39 μm x 39 μm x 100 μm, acquisition time = 1 h, 49 min.

DISCUSSION
RESULTS
The suitability of MEMRI to image the nervous system and other soft tissues of adult crayfish has previously been shown using different MEMRI procedures (Herberholz et al., 2004; Brinkley et al., 2005). These studies established that systemic injections of MnCl2 into the open circulatory system of crayfish significantly improved signal intensity in MRI images when compared to control animals that were not injected. Moreover, the time course of
manganese uptake into the brain and subsequent elimination from the brain was determined (Herberholz et al., 2004). In the only other published study using MEMRI in an invertebrate, Watanabe et al. (2006) investigated neural activity in both antennal lobes of Manduca sexta pupae after antennal transection on one side. Using T2-weighted imaging, they found a higher increase in SNR on the intact side as compared to the disrupted side indicating normal synaptic activity at pupal stage P13.

The current study supports previous reports that MEMRI is a highly successful technique to visualize the nervous system of live invertebrate animals. Moreover, we have demonstrated that MEMRI can be successfully used to label activity-dependent Mn2+ uptake into the brain and ventral nerve cord. Thus, MEMRI is well suited for both non-invasive neuroanatomical studies and functional studies of brain activity in various animals with small nervous systems. Since crayfish tolerate long imaging sessions in intense magnetic fields, MR images of high spatial resolution can be achieved. While overall spatial resolution cannot compete with conventional microscopic techniques, MEMRI is performed non-invasively in live animals and images can be generated within minutes to hours.

Although manganese is a strong T1 relaxation agent, our T2-weighted RARE images were characterized by high signal-to-noise. This is in agreement with previous studies and suggests that RARE pulse sequences may be well suited for anatomical studies using MEMRI or conventional MRI (Brinkley et al., 2005; Michaelis et al., 2005). T2-weighted images are also helpful for post-imaging data processing. They provide sufficient signal intensity across all brain areas and can guide ROI selection for quantitative measurements (e.g., grayscale values). In addition, detailed neuroanatomical structures can be visualized using 3D RARE scanning sequences which provide images that are not affected by positional changes of the animal inside the imaging tube and allow for better intra- and inter-individual comparisons.

For detection of localized Mn2+ uptake during functional MEMRI experiments, however, we found T1-weighted MSME imaging sequences to be most effective. The localized uptake of Mn2+ into one side of the brain after ipsilateral stimulation of the antenna II nerve was more clearly visible in T1-weighted MSME images, suggesting that this imaging sequence best reveals activity-dependent manganese uptake.

For analyzing grayscale values in the brain, we selected rectangular ROIs for each animal and used the same ROIs for both the stimulated and unstimulated sides of the brain. This allowed us to include all regions of the brain that are expected to be activated by the stimulation, and to compare the area to the exact same area on the unstimulated side. However, rectangular ROIs would sometimes include regions outside the brain that contained tissues of varying signal intensity, which slightly affected grayscale values. We explored an alternative method by hand-drawing localized ROIs around brain regions, and thereby excluding surrounding tissues. We found this challenging, however, as the brain outline was not always clearly detectable, and it was nearly impossible to use the same ROIs for both sides of the brain. Overall, using rectangular ROIs was a more conservative way to measure grayscale values in both halves of the brain, and yet revealed statistical differences between the stimulated and unstimulated regions.

Measurement of grayscale values showed differences between the two sides of the brain in all five stimulated animals; higher values were always found on the stimulated side indicating more Mn2+ uptake into the part of the brain that exhibited higher neural activity due to the stimulation. However, the differences between brain areas ipsilateral and contralateral to the electrically stimulated antenna II nerve varied across animals. This variation may be explained by differences in activity pattern of individual crayfish. We did not restrain the animals during stimulation and noticed that differences between the stimulated and unstimulated sides of the brain were less pronounced in crayfish that were more physically active during this period (e.g., some animals repeatedly tail-flipped in response to the stimulus while others did not). Thus, it seems possible that behaviorally more active animals experienced higher overall brain activity leading to an increase in non-localized Mn2+ uptake, which then masked the effects of antenna II stimulation. We are currently carrying out additional experiments to investigate this possibility.

Crayfish (and many other invertebrates) have large neurons, allowing MEMRI resolution to be sufficient for labeling of individual active cells. Our images of the third abdominal ganglion after repeated electrical stimulation of the LG circuit hint on this possibility. The bilateral pattern of highly localized uptake suggests that we imaged some of the large neurons that are part of this circuit (Zucker et al., 1971; Wine, 1984; Edwards et al., 1999). However, more experiments are needed to confirm this notion, and the localization of manganese labeled neurons must be carefully compared to existing morphological descriptions of this circuit before firm conclusions can be reached.

CHALLENGES

Very fast T1-weighted sequences such as fast low angle shot (FLASH) or echo planar imaging (EPI) can be used for manganese uptake studies. However, with our crayfish model these gradient echo based techniques yielded suboptimal results due to imaging artifacts caused by large air-filled areas inside the animal. The susceptibility differences between air and surrounding tissue results in additional field gradients on top of the imaging gradients which do not get refocused by the gradient echo and therefore result in marked signal loss and image distortion (Michaelis et al., 2005). At higher magnetic fields these effect are magnified, thus to minimize the effect of air-filled cavities on neuroanatomical images it is advisable to use imaging protocols based on spin echo techniques.

The signal intensity and maximum spatial resolution that can be obtained for an MR image are directly proportional to the magnetic field strength and gradient strengths. Higher field strengths are especially beneficial for studying smaller animals since their tomographic slices have lower spin density resulting in lower signal. The thickness of a slice imaged is further limited by the capability of the slice selective RF pulse generator in conjunction with the gradient field strength. Furthermore, for any 2D slice selected, the resolution is a function of FOV/matrix size, where the matrix size depends on the number of data points collected in the phase encoding and frequency encoding directions (the axes orthogonal to the selected slice). While the matrix size depends on the user input and should be based on the amount of time available for imaging (phase encoding steps increase acquisition time) the FOV...
is inversely correlated to the gradient strength. Hence, depending on the spatial resolution desired for the sample all the above factors must be taken into consideration for the choice of field and gradient strengths.

The injected amount of manganese and the temporal relationship between injection and the start of the imaging session is not critical for neuroanatomical studies. In fact, higher concentrations and amounts of manganese yield the best results because neural tissue is more intensely labeled and results in higher contrast between neural tissue and other surrounding tissues (Herberholz et al., 2004). Similar to previous reports, we found that animals used in our current study survived long periods inside the scanner (>24 h); however, the amount and concentration that can be tolerated by the animals must be newly determined for each species and each developmental stage and adjusted to size and/or weight of the sample. Moreover, not all studies will require Mn\(^{2+}\) as a contrast agent and conventional MRI can be used if contrast between different neural structures is high. This was demonstrated by Michaelis et al. (2005) using \(T_1\)- and \(T_2\)-weighted 3D MRI to describe morphological changes during metamorphosis in male pupae of M. sexta, although at a lower spatial resolution compared to our study.

In functional studies, temporal resolution of MEMRI images is associated with two factors; first, the time between manganese injection and the start of the imaging session, and secondly, the time needed to sample images at sufficient spatial resolution. Although manganese is quickly taken up from the bloodstream and slowly eliminated from the nervous tissue of crayfish (Herberholz et al., 2004), these factors can complicate unambiguous identification of stimulus-evoked neural activity. This is because our current knowledge predicts that all active neurons will take up manganese, and manganese ions are also transported across synapses from one neuron to the next albeit at a very slow rate. Thus, in order to identify a specific ensemble of neurons that is activated by the stimulation procedure, it is necessary to clearly distinguish its activity from other “unrelated” neural activity. Since manganese is injected into freely behaving animals outside the magnet, neural activity related to behavioral expressions as well as neural activity elicited by handling the animals during imaging preparation will cause manganese uptake into the activated neurons. This makes interpretation of the acquired images more difficult. One solution to this problem is injection of small amounts of Mn\(^{2+}\) that are only available in the bloodstream for a restricted time period, i.e., during and immediately following the stimulation of sensory pathways. Although this was attempted in our study, variation in mean grayscale values indicate that uptake into the brain and possible transport across brain areas has occurred at different times for different animals. In highly active animals, MnCl\(_2\) is probably circulated more quickly and uptake into the brain could start earlier than in less active animals. Thus, during the time of scanning (which we kept constant) the nervous tissue of each animal may have been exposed to different amounts of Mn\(^{2+}\).

**FUTURE APPLICATIONS**

Manganese-enhanced magnetic resonance imaging is a useful technique to investigate neuroanatomical features in vertebrates and invertebrates. The spatial resolution is high amongst non-invasive imaging techniques. Spatial resolution obtained with MEMRI may not compare to conventional microscopy, but it has the advantage of imaging live animals that can also be re-imaged after manganese has been eliminated from the neural tissue. This opens up new applications of MEMRI in a variety of different species, including recovery from injury and developmental studies when maturation of the nervous system can be tracked over several stages (Watanabe et al., 2006). Since RF inserts are available in different sizes, it may be possible to image animals from juvenile stage to adulthood (provided the typical size of the adult is smaller than the highest diameter insert).

Functional studies, if confirmed feasible by future experiments, will not have to be limited to protocols that measure neural activation by sensory stimulation. Invertebrates are characterized by large and accessible neurons, many of which are individually identified. It may therefore be possible to measure neural uptake into neurons and neural circuits that control specific patterns of behavior expressed in freely behaving animals. MEMRI holds the possibility of identifying CNS regions and perhaps single cells whose labeling changes as a result of behavioral changes. This would provide the necessary basis for further investigation of neural mechanisms using other techniques such as intracellular electrophysiology.

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