Safe use of subdermal needles for intraoperative monitoring with MRI

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OBJECTIVE The purpose of this study was to develop safe, site-specific procedures for placing and leaving subdermal needle leads for intraoperative monitoring (IOM) during intraoperative MRI procedures.

METHODS The authors tested a variety of standard subdermal needle electrodes designed and FDA-approved for IOM in the conventional operating room. Testing was used to determine the conditions necessary to avoid thermal injury and significant image artifacts with minimal disruption of IOM and MRI procedures. Phantom testing was performed with a fiber optic (lead) temperature monitoring system and was followed by testing of leads placed in a healthy volunteer. The volunteer testing used electrode placements typical of standard IOM cases, together with radiofrequency (RF) coil placement and imaging sequences routinely employed for these case types. Lead length was investigated to assess heating effects for electrodes placed within the RF coil.

RESULTS The authors found that conventional stainless steel (SS) and platinum/iridium (Pt/Ir) subdermal needles can be used safely without significant heating when placed outside the RF coil, and this accounts for the majority or entirety of electrode placements. When placed within the RF coil, Pt/Ir leads produced minimal image artifacts, while SS leads produced potentially significant artifacts. In phantom testing, significant heating was demonstrated in both SS and Pt/Ir leads placed within the RF coil, but only during high-resolution T2-weighted scanning. This problem was largely, but not completely, eliminated when leads were shortened to 25 cm. Human testing was unremarkable except for nonpainful heating detected in a few electrodes during thin-slice (1.5 mm) FLAIR scanning. Transient irritation (skin reddening along the needle tract) was noted at 2 of the electrodes with detectable heating.

CONCLUSIONS The authors were satisfied with the safety of their site-specific procedures and have begun with off-label use (following institutional review board approval and obtaining patient informed consent) of tested monitoring leads in cases that combine IOM and MRI. The authors recommend that all facilities perform their own site-specific testing of monitoring leads before proceeding with their routine use.

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Key words intraoperative monitoring; intraoperative MRI; MRI safety testing; MRI thermal injury; MRI artifact

Intraoperative monitoring (IOM) has been used in the operating room (OR) for more than 2 decades and provides a means of neurophysiological evaluation in surgical patients under general or local anesthesia who have significant neurological risk. A wide variety of monitoring modalities are available to assist surgeons in managing surgical risk through ongoing electrophysiological testing and mapping procedures on the brain, cranial nerves, spinal cord, and peripheral nerves. In recent years, a number of facilities have introduced an intraoperative MRI capability, which raises issues for the safe and effective use of standard IOM leads during MRI. After our medical center recently obtained an intraoperative MRI scanner, we were motivated to develop procedures for doing IOM safely and efficiently in the MRI environment. IOM typically employs subdermal needle electrodes inserted at...
número de localizaciones en el cuerpo para grabar y estimulación, que eleva serio la calidad de la imagen y consideraciones de seguridad. Los artefactos imagen pueden ser causados por el material compuesto de los electrodos, los cables, y el equipamiento.7 El riesgo principal de seguridad es la lesión térmica debido al electrodos durante la resonancia magnética (MRI) escáner, como un número de estudios han reportado lesions térmicas asociadas con MRI y otros tipos de cables (i.e., electrocardiograma y oxímetro de pulso leads).3,5,6 Desafortunadamente, en la práctica, MRI efectos térmicos son relativamente impredecibles y no se mueven de manera uniforme a través de diferentes instalaciones y condiciones operativas.3,13,14

Al momento de escritura de esta, no existen disponibles FDA- aprobado, MR-secure, o MR-conditional subdermal needle leads que se usan también para el IOM general. El "MR-secure" sólo aparece en la MRI seguridad información, mientras "MR-conditional" es más común, con etiquetado de elementos habiendo sido probado y aprobado para el escáner de resonancia magnetica específico debido a su configuración y personalidad cada MRI instalación y su equipamiento, creemos que la seguridad específica aprobado para MRI se debe realizar incluso cuando MR-conditional leads se hacen disponibles y cuando cambios en MRI equipamiento, secuencias de escaneo, o leads son contemplados.

Nuestro procedimiento estándar de procedimiento para realizar IOM con nuestro MRI escáner en su primer año de operación se realizó 1) el lugar de los cables después de inducir anestesia y realizar cualquier estudio preoperatorio MRI, 2) remover y hacer cuenta para nuestro monitoreo de los cables bajo la sonda RF anterior a intraoperatorio MRI, y, si es necesario, 3) reemplazar los cables monitoreo sin comprometer el campo estéril para continuar monitoreo. Otros centros en los EE.UU. y en el extranjero han reportado que se sigue el mismo (remover/replacement) abordaje para intraoperatorio MRI cuando realicen IOM. Sin embargo, removable/replacement significativamente extender el tiempo de cambio de los cables para intraoperatorio MRI, y algunos riesgos se involucran con la entrada de cables que se realizan bajo el equipo quirúrgico, especialmente si son cerca de o dentro del campo estéril. Muchos de los puntos de entrada no pueden ser visible una vez que están bajo el equipo quirúrgico, causando lead removable/replacement deben ser realizados por feel, que aumenta el riesgo de las agujas, inadvertido pullout de intravenosos catéteres o el monitoreo de otras anestesia líneas, contaminación del campo estéril, y potencial modificación del paciente’s original posicionamiento y padder.

Nuestra meta fue desarrollar prueba de seguridad, sitio-especifico procedimientos para colocar y dejar subdermal needle leads para intraoperatorio MRI estudios. Se debe notar que nuestro IOM equipamiento (amplificadores, estimuladores, cables, computarizado monitoreo sistema) no es MR compatible, y no deseamos contemplar la necesidad de monitorizar durante intraoperatorio MRI.

## Methods

### Leads and Imaging Procedures

Nuestro test se enfocó en 1) la seguridad en términos de la posibilidad de una lesión térmica debido a la electrodo, y 2) el análisis de la calidad de la imagen para evitar artefactos que provienen de cables MRI el campo de la imagen (FOV). Varios candidates platinum/iridium (Pt/Ir) y acero inoxidable (SS) subdermal needle leads fueron empleados para test de demostración (Tabla 1). Todos los cables subdermal leads fueron diseñados y aprobado por FDA para el uso del IOM en la regular OR, teniendo estándar DIN plug para conexión en el equipamiento MRI, fueron embalados estéteres, y estaban disponibles para compra. Adicional test fue realizada para evaluar la características de calor de SS cables de sujeción estables (VisiStat 35 W, Teleflex Medical Inc.) que son generalmente usados para secure cables en el cabello dentro de la línea del cabello.

A total of 12 separate MRI sessions were completed over the course of 2 months, including 6 phantom and 6 healthy volunteer tests. Scanning was performed in our intraoperative MRI scanner (IMRIS Inc./Skyra 3T, Siemens Medical Solutions, Inc.) using the same radiofrequency (RF) coil arrangement (2 intraoperative, 4-channel, receive-only phased-array flex coils, 1 upper and 1 lower for a total of 8 channels) and imaging sequences employed for intraoperative MRI. This included T1-weighted magnetization-prepared rapid gradient echo (MPRAGE), T2-weighted turbo spin echo (TSE), and T2-weighted fluid attenuated inversion recovery (FLAIR) imaging, which were run repeatedly while changing the lead orientation and position within and with respect to the RF coil. Both thin- and thick-slice (1.5 and 3 mm, respectively) FLAIR scans were investigated because electrode heating was noted during thin-slice FLAIR imaging when leads were placed within the RF coil (Tabla 2).

### Phantom Lead Testing

Nuestro test inicial evaluó subdermal needle electrodo plazas en un fantasma (honeydew melon approximating the size of a human head). Los electrodo se insertaron a una relativamente shallow angle under the melon skin to mimic the subdermal placement typically used for electroencephalography (EEG), electromyography (EMG), and stimulation leads and at various orientations within, at the edges of, and well outside the RF coil. Initial scanning was done with the leads (4 at a time given the limitation of 4 temperature measurement channels) at their manufactured length, using various schemes for configuring and routing the electrode tails out of the magnet. Lead length was investigated in some detail to assess the heating effects of MRI on leads placed within the RF coil.

### Table 1. Subdermal needle leads tested

<table>
<thead>
<tr>
<th>Description</th>
<th>Material</th>
<th>Dimensions (mm)</th>
<th>Tail (m)</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Straight needle</td>
<td>Pt/Ir</td>
<td>0.4 x 12</td>
<td>1.5</td>
<td>Technomed</td>
</tr>
<tr>
<td>Straight needle</td>
<td>SS</td>
<td>0.4 x 13</td>
<td>2.5</td>
<td>Rhythmlink International LLC</td>
</tr>
<tr>
<td>Straight needle pair</td>
<td>SS</td>
<td>0.4 x 13</td>
<td>2.0</td>
<td>Medtronic plc</td>
</tr>
<tr>
<td>Straight needle pair</td>
<td>SS</td>
<td>0.4 x 13</td>
<td>1.5</td>
<td>Rhythmlink International LLC</td>
</tr>
<tr>
<td>Corkscrew needle</td>
<td>SS</td>
<td>0.4 spiral</td>
<td>1.2</td>
<td>Medtronic plc</td>
</tr>
</tbody>
</table>
During phantom scanning, ongoing lead temperature measurement was performed using an MR-compatible, 4-channel, fiber optic (FO) temperature measurement system (Reflex, Neoptix Corp.). The FO sensors, which extended 5 m from the recorder (not MR compatible), recorded the temperature every second with a resolution of ± 0.1°C. Lead temperatures were measured with the FO sensors placed 1) under the melon skin parallel to the leads and separated by approximately 1 mm, and 2) in almost direct contact with the needle lead at its entry point through the melon skin. The second method was quickly adopted for most of the phantom testing, as it gave the most consistent and direct temperature measurements (Fig. 1).

**Lead Testing in a Healthy Volunteer**

A healthy volunteer (T.M.D.) was used to assess heating effects by placing leads to simulate typical IOM cases, including brain and upper and lower spine surgeries. Testing on a single individual was considered appropriate for the immediate aims of our site-specific safety testing, as opposed to a systematic research study intended to generalize findings to other facilities, which would have also required institutional review board approval. To assess lead temperature, we relied on the volunteer’s report of sensory changes during MRI and inspected the electrode sites after scanning. This was considered the most sensitive measurement technique under the circumstances. Using the FO temperature sensors on a healthy volunteer would be problematic from a practical standpoint, since the sensors would be difficult to sterilize and safely place in near contact with inserted needles without risking infection. How this might be facilitated in the future is considered in the Discussion below.

As with the phantom testing, multiple subdermal lead types from 3 different vendors were tested in the MRI scanner. Securement and placement techniques were also evaluated to identify potential safety or image quality issues. Except for the Pt/Ir leads, which were sutured in place with silk ties, securement and placement techniques were the same as those used in the regular OR. Additionally, strain-relieving loops were avoided. Leads outside the hairline were secured using surgical adhesive tape (Blenderm, 3M Healthcare) combined with liquid adhesive (Mastisol, Ferndale Labs). Leads within the hairline were similarly secured given that surgical staples and sutures could not be used without undue focal discomfort and consequent interference with sensory detection of electrode heating. In 1 case, a surgical staple was placed on the scalp.

**Results**

**Phantom Lead Testing**

As expected, the Pt/Ir leads created minimal imaging
artifacts, while the SS leads and surgical staples created significant localized artifacts up to 2 cm in diameter (Fig. 2). In the first 4 sessions, no significant temperature changes (> 4°C) were measured, and no clear inductive RF heating was observed with intentional looping and crossing of leads. In the fifth session, significant temperature changes were noted during T2-weighted scanning for 1 of the Pt/Ir and 1 of the SS electrodes, with maximum transient temperature changes of approximately 15°C in thin-slice FLAIR and 6°C in TSE sequences. This prompted immediate experimentation with lead lengths (through trimming of the electrode tails) to see if temperature increases could be reduced or eliminated. This line of investigation was prompted by reports of RF antenna-effect heating at particular lead lengths related to the MRI resonant wavelength. In fact, during T2-weighted imaging, heating was greatly diminished when leads were shortened to 40 cm and substantially eliminated when leads were shortened to 25 cm (Fig. 3).

Our summary assessment at this point was as follows: 1) leads of any type were safe in terms of heating when placed well outside the RF coil or subjected to T1 imaging only, 2) we needed to do further testing on the effects of lead length on heating during T2-weighted imaging, and 3) we would probably need to restrict lead length to prevent the heating of leads placed within the RF coil when performing thin-slice FLAIR and TSE scans. We also examined available information on lead length and focused on the 3 types of scalp leads with MR conditional FDA labeling (Table 3). These leads all have short tail lengths, ranging from 13 to 25 cm, and were packaged with removable MR-incompatible extensions of 200 to 250 cm. Based on this information, we decided to adopt 25 cm as the length we would pursue for leads placed within the RF coil, with the plan of using similar removable extensions to connect to IOM equipment, which would be disconnected and removed for intraoperative MRI. We should note that we had previously obtained evaluation samples of the above electrodes and concluded that none was appropriate for general IOM use. For subdermal needles with a reduced lead length of 25 cm, no additional significant temperature increases were measured in our phantom studies. At this point, we decided that it would be safe to test subdermal Pt/Ir and SS leads in a healthy volunteer.

Lead Testing in a Healthy Volunteer

The first MRI test was performed with leads placed on the foot and lower leg of the healthy volunteer, with his foot inside the RF coil. Full-length leads (1.5–2.5 m) were used, and no heating or skin irritation and/or injury was noted for any lead during MPRAGE, TSE, and thin-slice FLAIR imaging. Next, we performed extensive lead placements for 3 common types of surgery, followed by MRI while the leads were left in place. Based on results of the phantom tests, leads within the RF coil were shortened to 25 cm after MPRAGE imaging and prior to TSE or FLAIR imaging. As expected, MPRAGE imaging resulted in no significant heating of leads of any type or at any length when placed inside or outside the RF coil, and no heating was detected with any of the imaging modalities when leads were placed outside the RF coil. It should be emphasized that this accounts for the overwhelming majority of leads typically placed for IOM. And, as expected, imaging artifacts were insignificant for Pt/Ir leads placed within the RF coil but were potentially significant (1–2 cm diameter) for SS leads and surgical staples within the RF coil; therefore, one must choose between these electrode types on a case-by-case basis when placing leads within the RF coil. As detailed below, no differences were seen...
between the Pt/Ir and SS lead types in terms of their heating characteristics.

We first tested the lead setup for lumbar-sacral spine surgery requiring transcranial motor evoked potential (tc-MEP), somatosensory evoked potential (SSEP), and EMG monitoring (Fig. 4). Full-length (2 m), twisted-pair SS leads were used exclusively, with EMG lead pairs placed in the bilateral upper (first dorsal interosseous) and lower (adductor longus, vastus lateralis, tibialis anterior, gastrocnemius, abductor hallucis brevis) extremity muscles; EEG and/or SSEP leads placed at scalp locations FPz, CPz, CP3, and CP4; SSEP and train-of-four stimulating lead pairs placed over the bilateral ulnar and posterior tibial nerves; tc-MEP stimulating leads placed at scalp locations C1 and C2; and common (COM) electrodes placed on 1 upper and 1 lower extremity. Anal sphincter leads typically used in such cases were not applied, as these leads would be difficult to tolerate for a healthy awake volunteer. Since anal sphincter leads would often be within the RF coil in this type of case, short-tailed leads would probably need to be used if T2-weighted scanning was performed. The healthy volunteer was prone for MRI, just as he would be positioned for this type of surgery, and no heating effects or image distortions were noted for any type of MRI. This was not unexpected, as no leads were placed within the RF coil.

The lead setup for cervical spine surgery was similar to that for lumbar-sacral spine surgery, except that we used a different selection of EMG leads and a different placement scheme for the tc-MEP scalp leads. EMG leads were placed in the bilateral upper (trapezius, deltoid, biceps, extensor carpi radialis, first dorsal interosseous) and lower (adductor longus, tibialis anterior, abductor hallucis brevis) extremity muscles, and tc-MEP stimulating leads were placed at scalp locations C3 and C4 rather than C1 and C2. The remaining stimulating scalp EEG and COM leads were placed in the same locations as for lumbar-sacral spine surgery. Full-length SS twisted-pair leads were used exclusively, as none would be within the RF coil. The volunteer was prone for MRI, just as he would be positioned for this type of surgery, and no heating effects or image distortions were noted for any type of MRI. As before, these results were not surprising, since no leads were placed within the RF coil.

The third lead setup was for right-sided brain tumor surgery. EMG lead pairs were placed in the contralateral face (masseter, orbicularis oris), upper extremity (trapezius, deltoid, biceps, extensor carpi radialis, first dorsal interosseous, adductor digiti minimi, abductor pollicis brevis), and lower extremity (adductor longus, vastus lateralis, tibialis anterior, gastrocnemius abductor hallucis brevis) muscles. SSEP and train-of-four stimulating leads were applied to the contralateral wrist and ankle, a COM lead was placed on the shoulder, and scalp EEG leads were placed in areas FPz, CPz, CP3, and CP4. The tongue leads typically used in such cases were not applied, as these leads would have been difficult to tolerate for a healthy awake volunteer. Since tongue leads would be within the RF coil in this instance, short-tailed leads would probably need to be used if T2-weighted scanning was performed. Lead removal/replacement probably would be done pending appropriate safety testing. If T2-weighted scanning was performed, this uncertainty with regard to safety would generally apply to untested lead placements, including the tongue, soft palate, eye muscles, and anal sphincter. MPRAGE scanning was performed in 3 head positions (right side up, supine, left side up), and no heating was noted in any lead. The head and face leads were trimmed to roughly 25 cm (± 2 cm) prior to T2 FLAIR thin-slice scanning. Transient, nonpainful heating was clearly noted on the CP3 scalp electrode in the right-side-up position, on the Cpz electrode in the supine position, and on the CP3 and FPz electrodes in the left-side-up position. The heating was a pulsatile, pinprick sensation at the electrode site, correlated to the pulse sequence, with rapid extinction of the sensation between pulses. Inspection of the electrode sites after scanning revealed visible skin irritation at the FPz electrode site only, characterized as redness along the needle track (Fig. 5). This skin irritation was not persistent, did not require treatment, and was no longer visible a few days after the procedure. However, this irritation established that electrode heating was still possible with shortened leads and suggested the need for further tests focusing on scalp leads placed within the RF coil when T2 or other high-energy sequences were required.

The next scanning session used a larger array of scalp leads (12 positions including FPz, Fz, F3, Cz, C3, P3, Pz, and F4) and placed these leads exactly the same as for the cervical spine surgery. A healthy awake volunteer was positioned for this type of surgery, and no heating effects or image distortions were noted for any type of MRI. This was not unexpected, as no leads were placed within the RF coil. The volunteer was prone for MRI, just as he would be positioned for this type of surgery, and no heating effects or image distortions were noted for any type of MRI. As before, these results were not surprising, since no leads were placed within the RF coil.

The lead setup for lumbar-sacral spine surgery with intraoperative MRI was similar to that for lumbar-sacral spine surgery. Full-length SS twisted-pair leads were used exclusively, as none would be within the RF coil. In this case, all leads were SS with full-length tails, since no leads were placed within the RF coil.

**TABLE 3. Currently available, FDA-approved, MR-conditional scalp EEG leads**

<table>
<thead>
<tr>
<th>Electrode/Description</th>
<th>Material</th>
<th>Tail Length (cm)</th>
<th>Lead Extensions (cm)</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Press-on (3-prong subdermal w/ applicator)</td>
<td>Nickel titanium</td>
<td>25</td>
<td>250</td>
<td>Rhythmlink International LLC</td>
</tr>
<tr>
<td>MR-conditional cup (surface glue on, electrolyte gel)</td>
<td>Plastic, Ag/AgCl coating</td>
<td>24</td>
<td>150–250</td>
<td>Rhythmlink International LLC</td>
</tr>
<tr>
<td>SWE (subdermal wire w/ needle applicator)</td>
<td>Silver wire, AgCl coating</td>
<td>13–25</td>
<td>150–250</td>
<td>Ives EEG Solutions Inc.</td>
</tr>
</tbody>
</table>

**FIG. 4.** Images of healthy volunteer setup, with lead placements simulating a lumbar-sacral spine surgery with intraoperative MRI. In this case, all leads were SS with full-length tails, since no leads were placed within the RF coil.
A1, F4, C4, A2, P4) that were pretrimmed to precisely 25 cm and all placed within the RF coil. In this case, we focused on thin-slice and thick-slice (1.5 and 3 mm, respectively) FLAIR sequences obtained for the 3 head positions (supine, right side up, left side up), and all placed within the RF coil. In this case, we focused on thin-slice and thick-slice (1.5 and 3 mm, respectively) FLAIR sequences obtained for the 3 head positions (supine, right side up, left side up). No detectable heating or skin irritation was noted for any of the sequences or head positions.

After the aforementioned tests, we believed there was sufficient evidence to conclude that for FLAIR scanning, the tested leads would be safe if restricted to 25 cm when placed within the RF coil, although it remained possible that some heating and skin irritation could occur even with these precautions. After testing, we obtained institutional review board approval for off-label use of the tested leads at our facility with informed consent by the patient and have since used them without incident (no skin irritation evident at any electrode site) in 5 patients, including 2 who underwent cervical spine surgery and 3 who had surgery for brain tumor. For these patients, electrodes within the RF coil were either trimmed or removed/replaced when T2-weighted scanning was performed.

We next secured factory-made Pt/Ir and SS leads with 25-cm tails. This electrode tail length was considered a reasonable precaution considering 1) the mixed results of the prior 2 scanning sessions using scalp leads inside the RF coil (1 session demonstrating heating and the other not), and 2) the fact that these leads had DIN connectors as opposed to cut ends. For this test, a mix of Pt/Ir and SS leads were placed on the scalp (Fz, Cz, F4, C4, P4 for Pt/Ir; F3, C3, P3, Pz for SS), and 3 head positions were tested, including supine, right side up, and left side up, with a focus on FLAIR scanning. As before, no heating effects were noted for MPRAGE or TSE scanning. Transient, nonpainful heating was clearly noted at the F4 and P3 scalp electrodes in all 3 positions with thin-slice FLAIR scanning. The heat was a pulsatile, pinprick sensation that correlated to the pulse sequence, with rapid extinction of the sensation between pulses. This sensation was very similar in character to, but less intense than, that in the prior session in which there was detectable electrode heating. Trace sensations were still detectable for thick-slice FLAIR scanning. Inspection of the electrode sites after scanning revealed visible skin irritation at the C3 electrode site only, characterized as redness along the needle track. This skin irritation was not persistent, did not require treatment, and was no longer visible a few days after testing.

Discussion

The purpose of our site-specific safety testing was to establish in-house guidelines for the safe use of subdermal needle electrodes for IOM during intraoperative MRI. Establishing these guidelines was meant to obviate the removal and replacement of leads when monitoring is interrupted for an MR image. Before undertaking this study, we were encouraged by reports from other groups who described the successful use of subdermal needles in their facilities.4,9 We emphasize that our results do not generalize to other facilities, electrode types, or scanning sequences. We fully expect to do similar, repeated testing for any significant changes at our site, such as MRI hardware, scanning sequences, or lead types. We recommend similar testing at other facilities prior to the adoption of techniques for routine patient use.

The importance of keeping IOM in place during intraoperative MRI should be emphasized. Removal and replacement of needle electrodes from underneath sterile drapes poses a real risk of sterile field contamination. Electrode removal takes extra time given the difficulty in working underneath the sterile drape, and electrode replacement can disrupt careful patient positioning, leading to potential pressure sores from misplaced bolsters or padding. Because of this difficulty, a surgeon may continue tumor resection without IOM, which could increase operative morbidity.

A few lead types (conductive plastic cup, press-on, and silver hookwire electrodes, see below) have been developed for scalp EEG recording and are FDA labeled as MR conditional, but these leads are not appropriate or adaptable for general IOM use, which requires leads appropriate...
for EEG, EMG, and electrical stimulation at many body locations. Subdermal needle electrodes have proven most suitable for these combined purposes. A few institutions have reported safe and effective off-label use of relatively standard, FDA-approved subdermal needle types that are left in place for MRI at their facilities. The problem to date has been that those are site- and equipment-specific findings that do not generalize to other facilities. We adopted a strategy of performing our own site-specific testing of subdermal needle electrodes that were FDA approved for IOM use in the conventional OR, with a view toward their potential off-label use in the MRI scanner. Pending an MR-conditional listing by the FDA, informed consent will be used to apprise patients of the off-label use of unapproved leads that have been tested locally to assure ourselves and the institutional review board of the nonsignificant risks to the patient.

In a review of the literature, we found a clear lack of consensus and empirical evidence on the physics and specific mechanism of thermal injuries in the MRI scanner. This reflects the fact that the MRI system is a complex electromagnetic environment into which the patient is placed with multiple leads and subcutaneous lines, forming a complex network of possible conducting paths internal and external to the patient. The standard explanation for heating in the MRI scanner is the induction of high eddy or lead currents by changing RF magnetic fields according to Faraday’s law. Another, less commonly cited, explanation is the induction of resonant effects in tissue or leads by changing electric fields, typically referred to as “antenna effects.” There have been several demonstrations of these effects under test conditions, but the specific configurations under which heating occurs have not risen to the point of predictability based on theory, making empirical testing a practical necessity. As such, all MRI equipment, scanning sequences, leads, and placement procedures should be tested prior to routine use and then re-evaluated if any significant changes are made.

We acknowledge that there are limitations associated with phantom testing in a melon and the use of only 1 human volunteer. There are potential conductive differences between a melon and live human tissue with regard to heating. Live human tissue also exhibits potential protective mechanisms against overheating, such as sweating, that could help mitigate thermal injury. We believe our study was thorough and robust as regards the extent of testing in our melon and our human volunteer; however, we believe that further testing in other human volunteers could help validate our results and provide potential safety data prior to widespread patient use.

Our testing established that detectable and arguably significant electrode heating was possible in a narrow set of circumstances, even with the precaution of using shortened leads within the RF coil. This was seen with high-energy sequences such as thin-slice FLAIR. While we believe that the potential injuries associated with thin-slice FLAIR would probably be tolerable to patients if such scans were critical to intraoperative decision making, we plan to avoid using these sequences because of the potential risk to our patients. Our findings suggest that lower-energy scans, such as thick-slice FLAIR, should be considered as an alternative to thin-slice FLAIR when possible. Our findings also reinforce the need to maintain close scrutiny of electrode sites within the RF coil when T2 or other high-energy sequences are used.

Whether the heating effects noted herein rise to the level of a significant tissue injury is a matter for discussion and clinical judgment; these effects should be considered in relation to the conventional use of subdermal leads and securement staples, which produce minor puncture wounds and skin irritation that may be visible for a few days after surgery. In the few instances in which heating was detected in our testing, our volunteer reported that the sensation was not painful, and visible skin irritation was rarely produced. It is pertinent to note that investigations of temperature thresholds for dermal injury have shown that nonpainful heating of even extended duration does not produce significant skin injury. While the threshold between an acceptable “minor” injury compared with an unacceptable “major” injury may vary from person to person, we believe that limiting imaging to lower-energy MR scans will keep potential thermal injuries at a level similar to the puncture wounds and skin irritation we typically encounter from traditional electrode placement.

A key breakthrough would be the development of a principled model that could predict and mitigate heating effects in the MRI scanner. This would help us and others to overcome the need for empirical testing and to devise more general and site-specific safety procedures. Simple, theoretical heating mechanisms such as Faraday induction and resonant antenna effects are hard to apply in practice. Improved measurement technologies capable of sensing lead temperature or induced currents would also be helpful, and there are a number of feasible options. One possibility is the development of a lead with a port into which an FO temperature probe could be inserted to make thermal contact with the lead without contaminating the subdermal needle. Another possibility is the use of infrared imaging technology that would allow visualization and measurement of the temperature distribution on the scalp or body surface, including the electrode sites. A third possibility is the use of MR-compatible, high-frequency electrical measurements (voltage or currents carried by the lead wires), which might be obtainable from unterminated DIN plugs on each lead. This latter possibility would perhaps allow detection of resonant or other high-current conditions and facilitate the introduction and testing of countermeasures, such as the use of additional components aimed at damping resonant oscillations or dangerous induced currents.

Conclusions

First, conventional SS subdermal IOM needles with standard-length (1.5–2.5 m) tails can be safely used outside the RF coil without significant heating in the tested scan types (MPRAGE, TSE, and FLAIR). In most IOM and MRI cases, this use of IOM needles outside the RF coil will account for the majority or entirety of electrode placements. Second, in phantom testing, significant heating of some electrodes was demonstrated for standard-length (1.5–2.5 m) SS and Pt/Ir leads placed within the RF coil, but only during T2-weighted scanning (FLAIR >
TSE), with no heating noted during T1-weighted scanning. SS and Pt/Ir electrodes appeared equally likely to exhibit heating, and no clear pattern of which electrodes would heat was evident. The maximum recorded temperature increase in phantom testing was 15°C in a Pt/Ir electrode. In addition, phantom testing of leads within the RF coil during T2-weighted scanning revealed the elimination of significant heating when leads were shortened to 25 cm. Thus, we decided to use short-tailed leads for placements within the RF coil when T2-weighted scanning was required. Third, imaging artifacts were minimal for Pt/Ir electrodes but were potentially significant (1–2 cm diameter) for SS electrodes within the RF coil and image FOV. Depending on the clinical circumstances, a mix of Pt/Ir and SS leads could be used, e.g., placement of Pt/Ir leads over the left hemisphere and SS leads over the right hemisphere in a patient with a left-sided tumor. On a practical note, Pt/Ir leads cost approximately 5 times as much as SS leads. Fourth, human testing was unremarkable except for nonpainful heating that was detected during thin-slice FLAIR scanning while using short-tailed 25-cm leads within the RF coil. This heating occurred sporadically at a small number (2–3) of Pt/Ir or SS electrodes in 2 of 3 sessions. Transient irritation (visible skin reddening along the needle tract) was noted at 2 of the electrodes with detectable heating. Heating became barely detectable when thick-slice FLAIR scanning was performed.

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References


Disclosures

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Author Contributions

Conception and design: Bauer, Darcey, Kobylarz, Roberts. Acquisition of data: Bauer, Darcey, Krauss, Pearl, Roberts. Analysis and interpretation of data: Darcey. Drafting the article: Darcey. Critically revising the article: Bauer, Kobylarz, Pearl, Ferri, Roberts. Reviewed submitted version of manuscript: Bauer, Darcey, Krauss, Pearl, Roberts. Administrative/technical/material support: Pearl, Ferri. Study supervision: Darcey.

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