Setting Up Workflow of an Intraoperative MRI Unit: A Single-Centre Experience of First 53 Cases

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Introduction

Intraoperative magnetic resonance imaging (iMRI) technology is a landmark development in this era of image-guided neurosurgery. The important advantages of iMRI are that it provides real-time imaging, locates abnormalities in case of brain shift, distinguishes abnormal brain tissue from normal, and guides complete and precise resection of tumor.
The iMRI setups can be of three types. First type comprises portable iMRI device which moves into the OR for imaging. In the second type, also called “nearby iMRI” suite, the MRI magnet is placed adjacent to the OR and patient is transported to the MRI suite for imaging. In the third type of iMRI setup, the MRI scanner is installed within the OR, and the MRI table can itself be used as the operating table.

Our setup is a nearby OR type, 3T-iMRI with the Siemens Skyra magnet located adjacent to the standard OR. This observational study of our first 53 cases during the first 3 months was performed to assess our learning curve of optimal resource utilization and conduct of cases using iMRI for the duration of October to December 2017. After installing the iMRI suite, mock drills were performed for 2 weeks to develop a protocol for shifting the patient to the iMRI, performing the scan, and shifting back to the operating room. Healthy volunteers were made to lie down on the operating table, and the maneuver of transporting the volunteer to the MRI suite and back was performed multiple times. The sequence of entry into the MRI suite was changed three times until the most comfortable sequence was finalized. All the personnel including scrub nurses and OR and MRI technicians who would be part of the team for iMRI were formally trained in didactic and practical session formats regarding the precautions and protocols to be followed. Warning signs and checklists were put on the entry door leading from OR to the MRI suite stating the MRI magnet would always be on and list of items prohibited inside the MRI suite at any time. Training was also conducted in categorizing all possible equipment, which could be present around and within the iMRI suite as MRI safe, unsafe, and conditional. A 5-Gauss line indicating strong magnetic field was marked inside the MRI suite. After the completion of training, actual surgical cases were started.

The surgeries were performed on a special operating table designed with an in-built MRI compatible sliding board for moving the patient on to the MRI gurney intraoperatively. MRI coil was placed beneath the patient’s head at the area of interest. Portable MRI compatible monitors were attached, and special MRI compatible carbon lead electrocardiogram (ECG) electrodes with braided wires were used. In the cases requiring evoked potential/electromyography monitoring, the subdermal needle/corkscrew electrodes used were “MR-conditional.” Darcey et al. tested the safety of subdermal needle and corkscrew electrodes in healthy volunteers and reported nonpainful heating and transient irritation in two and one site, respectively, of the 12 sites placed. They concluded that the MR-conditional electrodes can be used safely with iMRI. After induction of anesthesia, all skin to skin contact interfaces such as finger web spaces and perineum were padded with cotton pledges to prevent moisture accumulation and heating during MRI, which can cause burns. The patients were then positioned for surgery. MRI compatible head pins and navigation system were used while positioning (Fig. 1). After final positioning, a bore gauge was used to mimic the circumference of space available within the MRI magnet to confirm that the patient would slide smoothly in and out of the magnet, (Fig. 1) especially in patients positioned prone and lateral. After this step, the surgeon was asked to proceed with the surgery.

When the surgeon felt the need for an iMRI scan, the MRI technician and radiologist were alerted at least 30 minutes prior to the expected time for beginning the scan. The MRI suite was cleaned and sterilized during this time. The patient and surgical site were packed and covered with sterile dressing without closing dura so as to allow residual tumor resection after iMRI (Fig. 2a). Long intravenous (IV) line extensions (200 cm) and long breathing circuit extensions (1.5 m × 2 nos.) were connected to facilitate movement during MRI. MRI compatible infusion pump casing to carry infusion pumps during MRI was kept ready. Standard monitoring was continued with a portable TESLA MRI-compatible monitor for continuous monitoring of ECG, peripheral capillary oxygen saturation (SpO₂), end-tidal carbon dioxide (EtCO₂), and arterial blood pressure. A checklist for the patient and personnel going into the MRI suite was prepared during our training sessions to ensure patient, personnel, and equipment safety (Fig. 3). The checklist was read and confirmed by the anesthesiologist every time before moving the patient to the MRI.
suite. On completion of the checklist, MRI compatible gurney was brought in, docked with the operating table; and the patient was slid on to the gurney (Fig. 2a). The MRI compatible anesthesia workstation (GE Aestiva MRI; Datex Ohmeda Inc., Madison, Wisconsin, United States) in the MRI suite was checked and kept ready before moving the patient to the MRI suite (Fig. 2b). The breathing circuit was disconnected from the anesthesia machine in the operation theater and connected to the MRI compatible anesthesia workstation in the MRI suite. MRI compatible laryngoscope was checked and kept ready. Patient was taken into the MRI suite and during MRI scan, monitoring of vital parameters of the patient was continued, and a uniform depth of anesthesia was maintained (Fig. 2b). On completion of MRI, the MRI gurney with patient was brought back, re-docked to the operation table, and patient was moved on to the operation table. Any residual tumor detected on iMRI scan was removed or closure was performed after achievement of hemostasis.

Every time an iMRI was conducted, we recorded data regarding any mistakes, events, or complications while shifting the patient to the scan and back and during the scan, time required to move the patient into and out of the MRI suite, duration of the MRI scan, and number of personnel required to perform the iMRI scan. We also noted the number of patients in whom residue was detected and subsequently re-surgery for further resection was performed.

The data tabulation and calculation of descriptive statistics was conducted using Microsoft Excel 2010. Interval data were expressed as mean ± standard deviation, and categorical data were expressed as frequencies and percentages.

Results

This observational study was conducted over 3 months from October 2017 to December 2017 involving the first 53 cases who underwent neurosurgery with iMRI at our institute. As this study included an intraoperative noninvasive diagnostic procedure and the objectives of interest pertained to conduct of the procedure itself, the institute’s ethics committee waived off the need for a formal informed consent.

Table 1 shows the demographic data of the patients along with the diagnosis and position of the patient. Of the 53 cases that underwent iMRI, 51 were for tumor resection, and the rest 2 cases were patients of Parkinson’s disease for confirming position of deep brain stimulator (DBS) electrodes in the subthalamic nucleus. The most common surgeries for iMRI usage were transnasal trans-sphenoidal resection of pituitary tumors (26/53; 49.03%) and cerebellopontine angle tumor resections (12/53; 22.64%). Other surgeries in decreasing order of frequency were supratentorial tumor, posterior

Fig. 2 (A) MRI gurney docked with OT table before the patient is packed in a sterile fashion and OT tabletop slid on to the MRI gurney. (B) Patient on the MRI gantry with upper coil placed on top of the area of interest for scan. MRI compatible anesthesia workstation, portable monitor, and infusion pump casing can be seen. MRI, magnetic resonance imaging; OT, operation theater.
CHECKLIST FOR INTRA-OP MRI:

(To be Completed mandatorily prior to Wheeling-in; Please Tick on the boxes on checklist completion)

Name:  
Age:  Sex:  IP Number:  

MRI TIMING:  AM/PM TO  AM/PM

- MRI Compatible ECG electrodes:
- Coil placed
- OT Table levelled, stored position, Hemostasis secured at levelled position
- No metallic ornaments, Implants, Heart valves, Pacemakers in patient:
- Ear plugs for patient:
- Arms insulated from body, skin to skin padding
- No sand bags, metal objects in sheets, sisters’ instruments checklist done:
- OT table sheets tucked under to help sliding:
- Temperature removed:
- Urine Bag free:
- Warmer disconnected:
- Antibiotic repeated:
- MRI Technician informed:
- Cautery pad, RF pad, EM navigation removed
- Emergency drugs box sent in:
- Lumbar Drain clamped and secured:
- Monitoring Electrodes: (Entropy/Depth of anaesthesia) removed:
- OT personnel moving into the MRI Suite screened for metallic objects:
- MRI Compatible Monitor Ready:
  - Pulse oximeter:
  - EtCO₂:
  - ECG:
  - NIBP:
  - IBP:
- Anaesthesia Machine in MRI checked and ready:
- Infusion pump casing:
- Long breathing circuit(Double):
- Long IV line extension (200 ml):
- Long infusion extension (200 ml):
- AMBU, Bains circuit ready:
- Compatible Laryngoscope Ready:

VENTILATOR PARAMETERS:

Ventilation Mode:  

<table>
<thead>
<tr>
<th>TV</th>
<th>RR</th>
<th>FiO₂</th>
<th>PEEP</th>
<th>SEVOFLURANE</th>
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</tbody>
</table>

INFUSIONS:

1. ____________________ @ ______ml/Hr.
2. ____________________ @ ______ml/Hr.
3. ____________________ @ ______ml/Hr.

While shifting:

Urine Bag Free, Lines Free, Circuit Free

INFUSION PUMP WITH  MRI COMPATIBLE  PATIENT

PROTECTIVE CASING  MONITOR

Fig. 3  Checklist of our institute for patient, OT personnel and equipment safety to be performed before shifting for iMRI scan by the anesthesiologist. AMBU, artificial manual breathing unit; ECG, electrocardiogram; EM, electromagnetic; EtCO₂, end-tidal carbon dioxide; FiO₂, fraction of inspired oxygen; IBP, invasive blood pressure; iMRI, intraoperative magnetic resonance imaging; MRI, magnetic resonance imaging; NIBP, noninvasive blood pressure; OT, operation theater; PEEP, positive end-expiratory pressure; RF, radiofrequency; RR, respiratory rate; TV, tidal volume
fossa tumors, and arteriovenous malformation resections. A special mention is its use during awake craniotomy for intracranial tumor resection.

Out of the 51 tumor cases, residue was detected in 28 (54.90%) cases and no residue in 23 (45.09%) cases. Of the cases where residual tumor was detected on iMRI, re-surgery for further resection to achieve maximal or total resection was performed in 21/28 (75%) cases, whereas in 7 cases the residual tumor was not chased as it was very close to a blood vessel or a critical eloquent structure; thus, subtotal excision was performed (Table 2). The image quality of the scans was graded in a subjective manner on a 3-point scale based on the operating surgeon; in 45/53 (84%) cases, it was graded as excellent and provided vital information during performance of surgery (Table 2). In eight cases, surgeons were not satisfied with the image quality of iMRI scan. Image quality was rated poor in two cases in prone position and average in six cases with intraoperative neurophysiological monitoring (somatosensory and motor evoked potentials) due to the corkscrew scalp electrodes causing artifacts.

We were able to cut down on the number of personnel required to shift patients in and out of iMRI after multiple iterations of the procedure. After initial cases, only four personnel (one anesthesia consultant, one anesthesia resident, one anesthesia technician, and one MRI technician) were required to shift the patients and perform the scan. Total duration of MRI scan time was reduced after initial cases as the number of requisite sequences were reduced as per need. The mean duration of time for shifting patients for the iMRI, mean scan duration, and time required for shifting back are tabulated in Table 2. The shifting times to and from the iMRI suite reduced after first few trials (Fig. 4A and B). We also analyzed and compared the shifting time and scan duration in terms of first 26 and next 27 cases (Table 2). Although both the shifting and scan duration were reduced in the last 27 cases, the shifting duration significantly reduced (p = 0.014). We also prepared a thorough checklist for ensuring patient, personnel, and equipment safety during patient transport as well as during the scan. This checklist was changed five times as and when issues were identified.

We also encountered complications during these cases. One patient had burns in prone position where the lower coil was touching the patient’s nose, which was probably due to moisture collection. After this, we modified our technique in prone position and placed the lower coil just before moving into MRI scan to prevent blood and moisture collection. One patient had a swelling at the intravenous catheter site, after which we made it a protocol to make loops of IV line tubing to prevent cannula displacement due to tension on the IV line. Circuit got disconnected in one patient with a brief increase in EtCO₂ while moving into the scan, and corrective measures were taken by adding a step of “tightening of circuit connections before shifting” to the shifting workflow, to prevent recurrence in further cases. We also noted logistical issues during each step of iMRI and planned for corrective measures so that they would not recur in future cases. These were then incorporated while preparing the checklist.

### Discussion

Though intraoperative MRI provides vital information to the neurosurgeons in planning and performing neurosurgeries with precision, it poses a lot of challenges for the anesthesiologist team. Safety of the patient, OT personnel, and equipment are core issues during such diagnostics. With the help of our training sessions and data collection, we were able to develop an institutional protocol for effective resource utilization in terms of time taken and personnel required. The time taken and the number of personnel reduced sequentially over the course of the first few scans, as the technique was improved upon iteratively.

Challenges faced by the neuroanesthesia team are centered around moving the patient intraoperatively on head pins, monitoring and maintaining hemodynamics during the shifting procedure, and maintaining adequate anesthesia depth and adequate ventilation during the movement of patient into the MRI suite and back. Issues about MRI compatibility of equipment. Training of staff involved, and other OT personnel should be conducted and coordinated by the anesthesiologist who should be given the role of team leader. Team work and inter-departmental communication is key to the success of any iMRI unit.

Institutional protocols and checklists should be prepared and strictly adhered to for each case. Continuous data recording and auditing helped us reduce mistakes, personnel requirement, and also time taken to conduct the iMRI. Learning points from our experience in this regard to ensure smooth functioning of the setup have been summarized (Table 3).

| Table 1 Patient demographics, diagnosis, and intraoperative position of the patient |
|---------------------------------|---------------------------------|
| **Age (mean ± SD) years** | 50.26 ± 15.32 |
| **Sex (male:female)** | 28:25 |
| **Diagnosis** | **Pituitary tumors** 26/53 (49.03%) |
| | **Cerebellopontine angle tumors** 12/53 (22.64%) |
| | **Posterior fossa tumors** 3/53 (5.66%) |
| | **Supra tentorial tumors** 8/53 (15.09%) |
| | **Arteriovenous malformations** 2/53 (3.77%) |
| | **Parkinson’s disease (for deep brain stimulation)** 2/53 (3.77%) |
| **Position** | **Supine** 38 |
| | **Lateral** 12 |
| | **Prone** 3 |
| | **Awake craniotomies** 4 |

Note: The total number of procedures conducted is 53.
Table 2  Descriptive statistics of all collected data

<table>
<thead>
<tr>
<th>Description</th>
<th>Mean ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time to start shifting till time to start scan (mean ± SD) min</td>
<td>7.26 ± 6.50</td>
</tr>
<tr>
<td>Duration of scan (mean ± SD) min</td>
<td>26.0 ± 9.09</td>
</tr>
<tr>
<td>Duration to shift patient back from MRI suite to OR (mean ± SD) min</td>
<td>4.14 ± 2.96</td>
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<tr>
<td>Total duration including shifting in and out of MRI suite and scan time (mean ± SD) min</td>
<td>37.40 ± 13.50</td>
</tr>
<tr>
<td>Total duration required for shifting the patient in and out of MRI suite (mean ± SD) min</td>
<td>11.40 ± 9.01</td>
</tr>
<tr>
<td>Shifting duration: first 26 cases (mean ± SD) min</td>
<td>14.23 ± 11.27</td>
</tr>
<tr>
<td>Last 27 cases (mean ± SD) min</td>
<td>8.15 ± 3.80</td>
</tr>
</tbody>
</table>

*p-Value: 0.014

Total duration—first 26 cases (mean ± SD) min: 38.5 ± 16.26
p-Value: 0.39

Total number of personnel required (median) 4.00

Residue

<table>
<thead>
<tr>
<th>Status</th>
<th>Count</th>
<th>Percentage</th>
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</thead>
<tbody>
<tr>
<td>Yes</td>
<td>28/51</td>
<td>(54.90%)</td>
</tr>
<tr>
<td>No</td>
<td>23/51</td>
<td>(45.09%)</td>
</tr>
</tbody>
</table>

Re–surgery for residue 21/28 (75%)

Correct position of DBS electrode 2/2

Image quality

<table>
<thead>
<tr>
<th>Quality</th>
<th>Count</th>
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</thead>
<tbody>
<tr>
<td>Excellent</td>
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<tr>
<td>Average</td>
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<td>Poor</td>
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Complications

<table>
<thead>
<tr>
<th>Issue</th>
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</thead>
<tbody>
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<td>Burns</td>
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<tr>
<td>Circuit disconnection during shifting and rise in EtCO₂,</td>
<td>1</td>
</tr>
<tr>
<td>Hemodynamic disturbances</td>
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<tr>
<td>Hypertension and tachycardia</td>
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<tr>
<td>Hypotension</td>
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<tr>
<td>Hematoma detected on iMRI</td>
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<tr>
<td>IV line displaced with local swelling</td>
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<tr>
<td>Small technical and corrective issues</td>
<td>18</td>
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<tr>
<td>Issue during OT table docking</td>
<td>5</td>
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<tr>
<td>During shifting into scan</td>
<td>4</td>
</tr>
<tr>
<td>Docking into MRI</td>
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<tr>
<td>During MRI scan</td>
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<tr>
<td>Shifting back to OT and re–docking</td>
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<tr>
<td>Charging of MRI compatible monitor:</td>
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</table>

Abbreviations: DBS, deep brain stimulator; EtCO₂, end–tidal carbon dioxide; iMRI, intraoperative magnetic resonance imaging; MRI, magnetic resonance imaging; OR, operation room; OT, operation theater; SD, standard deviation.

Conclusion

While setting up any new interventional or investigative modality has its challenges, iMRI set up is a multi-departmental effort with significant issues pertaining to resource utilization and patient and personnel safety. Our experience shows that it takes multiple iterations of the procedure, coupled with initial training sessions and meticulous data collection and auditing to smoothen out the workflow.
A Single-Centre Experience of Setting Up of a New Intraoperative Magnetic Resonance Imaging Unit and Analysis of Resource Utilization  Manohar et al.

Table 3 Learning points from our experience to ensure smooth functioning during setup of the new system

1. Categorize all equipment: MRI safe, unsafe, and conditional
2. Mock drills with healthy volunteers to develop initial protocol for shifting patients into iMRI and back to operating table.
3. Intensive training of all OT personnel—nursing staff, OR, and MRI technicians
4. Warning signs/checklists posted at strategic points
5. MRI compatible workstation/monitors/infusion pumps—checked prior to use
6. Burns prevention measures: cotton pledges placement at skin to skin contact interfaces
7. Long intravenous extensions and breathing circuit with firm attachment and adequate slack to prevent disconnections
8. Strict adherence to detailed checklist customized as per institute practices and resources

Abbreviations: iMRI, intraoperative magnetic resonance imaging; MRI, magnetic resonance imaging; OR, operation room; OT, operation theater.

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None.

Conflict of Interest
None.

References