COMPUTER-ASSISTED INTRA-OPERATIVE MAGNETIC RESONANCE IMAGING MONITORING OF INTERSTITIAL LASER THERAPY IN THE BRAIN: A CASE REPORT

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ABSTRACT

Hardware and software for a customized system to use magnetic resonance imaging (MRI) to noninvasively monitor laser-induced interstitial thermal therapy of brain tumors are reported. An open-configuration interventional MRI unit was used to guide optical fiber placement and monitor the deposition of laser energy into the targeted lesion. T1-weighted fast spin echo and gradient echo images were used to monitor the laser tissue interaction. The images were transferred from the MRI scanner to a customized research workstation and were processed intraoperatively. Newly developed software enabled rapid (27–221 ms) availability of calculated images. A case report is given showing images which reveal the laser–tissue interaction. The system design is feasible for on-line monitoring of interstitial laser therapy. © 1998 Society of Photo-Optical Instrumentation Engineers. [S1083-3668(98)01503-2]

Keywords magnetic resonance imaging; interventional MRI; computer assisted surgery; CAS; surgery; minimally invasive surgery.

1 INTRODUCTION

Laser-induced interstitial thermal therapy (LITT) is a minimally invasive surgical technique for the treatment of solid tumors. LITT has long been a topic of research; studies have focused on application of the technique in a variety of normal tissue types and tumors in vivo and in vitro.1—10 The technique involves the percutaneous introduction of an optical fiber through a needle; light delivered to the tissue is absorbed proximal to the tip and the heat generated creates a localized coagulative necrosis. As the device is interstitial, the laser–tissue interaction occurs remotely from the operator/surgeon and is not visible as would be the case in an open surgical procedure. Monitoring and control of the treatment is needed. One solution has been through the use of radiologic imaging techniques such as magnetic resonance imaging (MRI) and ultrasound (US) which can provide images of the interaction.2,7,11—24

Radiologic imaging techniques offer: (i) accurate image-guided placement of the optical fiber(s) into the target; (ii) intra-operative imaging to monitor tissue changes as they occur; and (iii) feedback (i.e., images of acute thermal effects) by which dosimetry can be optimized by additional treatment or a repositioning of the fiber. These provide control of the location and spatial extent of the thermally affected area (i.e., “treatment site,” “induced lesion”). MRI has found considerable acceptance and the clinical experience (of image-guided LITT in the brain, head and neck, liver, and spine) at various centers has been reported.25—34

It is important that MRI has excellent resolution of soft tissue but it is as important that various “types” of MRI are temperature sensitive (i.e., the signal intensity on the image is a function of temperature).35 Types of MR imaging can be characterized by the “pulse sequence” by which the images are obtained. MRI pulse sequences such as those providing T1-weighted (T1w) fast spin echo (FSE), diffusion-weighted, and chemical-shift imaging have been shown to be temperature sensitive.11,13,36—38

MRI can provide a noninvasive means by which to monitor LITT. MRI sequences now provide images on a time scale which is suitable for LITT, providing near real-time monitoring. As changes in technology enable MRI-guided LITT to be implemented on patients, the information in the images must be optimally displayed so as to best appreciate the laser effect and to speed procedures. During...
LITT changes in the images’ signal intensity around the laser fiber’s tip ranges from strong to subtle. This article presents the implementation of a research computer workstation to assist in the manipulation of image data to provide needed feedback as to the laser effect in a clinical setting. We report on one patient treated by LITT in the brain under both a T1w FSE and a water proton chemical shift imaging sequence for on-line monitoring of the intra-operative thermal effect.

2 MATERIALS AND METHODS

2.1 MR IMAGING

MR images were acquired using an “open configuration” interventional MRI (iMRI) unit (Signa SP, General Electric Medical Systems, Milwaukee, WI).39 (See Figure 1.) The 0.5 T unit consists of upright coils spaced 0.56 m apart. The space allows a surgeon access to the patient while the anatomy is within the imaging volume (Figure 2). The MR images are displayed to the surgeon on liquid crystal displays (model LQ6NC01, Sharp Electronics, Rahway, NJ) located above the surgical field. The iMRI system is well suited to provide good access to the patient while drawing trajectories to targeted tissue, inserting probes/needles under image guidance, and viewing images of the anatomy intra-operatively—all while the concerns of anesthesia, nursing, and sterility are met. To date, this system has been successfully applied to biopsies, tumor resections, and endoscopic surgeries.40–46

Two modes of scanning have been used: (1) conventional T1w FSE and (2) fast two-dimensional (2D) spoiled gradient recalled echo (2D-FSPGR). The T1w FSE has been cited to have a sensitivity of its signal intensity of 0.48%/°C (FSE, TR=300 ms, TE=12 ms) in a gel phantom.36 The 2D-FSPGR, our water proton chemical shift sequence, has been reported to have a temperature sensitivity of −0.0135 ppm/°C.37

2.2 LASER DEVICE AND SETUP

The laser device (Sharplase 60, Sharplan Lasers, Allendale, NJ) emitted a continuous Nd:YAG laser at 1064 nm (nanometers). The laser light was transmitted into the procedure room through a long optical fiber passing through a port in the operating suite’s magnet-shielded access panel. The long fiber passed the light into a connecting box (1-to-3 optical beamsplitter; Sharplan Lasers, Allendale, NJ). Up to three delivery fibers can be attached to the connecting box. The delivery fibers, which carry the light into the tissue, are bare, sterilized 600 μm fibers. The laser irradiation was performed at two positions in the tumor. The delivery fiber output was confirmed pre-operatively with an external power meter to be 4 W.

Positioning of the fibers in tissue was achieved using interactive scan plane definition by which an oblique image plane is determined using an optical scan-plane locator (Flashpoint, IGT Inc., Boulder, CO). This locator has three light-emitting diodes which are tracked by three charge coupled device (CCD) cameras attached to a gantry above the interventional field. The oblique scan plane can match an arbitrary plane for the trajectory of the needle into the tumor.

2.3 RESEARCH WORKSTATION

The iMRI unit is equipped with a dedicated workstation (Sun 4 model 670, Sun Microsystems, Mountain View, CA) by which the unit is run. We installed a second workstation (SPARCstation 20TZX model HS21, Sun Microsystems, Mountain View, CA) adjacent to the iMRI workstation. Our image processing computations were performed on this research workstation, since the hardware resources...
of the MR workstation must be reserved for the image acquisition. The research unit was suitably capable of handling fast graphics manipulation and network communication needs. The research workstation had two separate display outputs each with its own graphics card: (1) a standard graphics card (TurboGX, Sun Microsystems, Mountain View, CA) to control the graphical user interface (Figure 3) which was connected to a monitor display (20 in. color monitor, Sun Microsystems, Mountain View, CA) placed in the iMRI control area, and (2) a card (TurboZX, Sun Microsystems, Mountain View, CA) which supports graphics acceleration and feeds the NTSC video signal to the two displays over the surgical field.

### 2.4 NETWORK

The iMRI unit, its workstation, and the research workstation used a standard TCP/IP protocol interface to establish network connection. Hosts were also connected to the hospital’s network which connects existing research facilities, such as mass storage of pre-operative data (currently more than 180 GByte), connections to conventional MR/CT scanners and high end computing resources. However, the system presented in this article can operate independently, without access to these research facilities.

The network was based on switched Ethernet with a faster backbone, using high bandwidth asynchronous transfer mode (ATM). ATM bridges the switched Ethernet network, to which the research workstation belongs, and a second switched Ethernet based network, which contains the iMRI workstation. The data transfer speed is 10 Mbps in the switched Ethernet network and 155 Mbps in ATM. Therefore, each machine can achieve a maximum transfer rate of 10 Mbps. Typically, the data transfer time is 0.72 s for an image of $256 \times 256 \times 16$ bits.

### 2.5 SOFTWARE DEVELOPMENT

Figure 4 presents a schematic of the overall control flow. Server/client software was developed and installed in the iMRI workstation and the research workstation, respectively. This software distributed tasks by establishing a network communication by remote procedure control (RPC) and transferred commands and images between them.

The iMRI server software is used to obtain image data from the scanner and send it to the research workstation. On request from the research workstation, the iMRI workstation allows access to its image buffer for transfer of the most recently acquired image to the research workstation. The image-to-buffer routine works independently of the server software and continuously refreshes its contents.
when the real-time scan is performed. This buffer is also shared with real-time image viewer and controller, which is installed in the iMRI workstation as the default interface.

The research workstation receives real-time images and processes them to provide a difference image to serve as a guide to the LITT procedure. This difference image can be routed for display to display screens above the surgical field.

Since the research workstation is dedicated specifically for 2D computer graphics, image processing, and its own customized user interface, its hardware is configured to maximize the performance of these functions. The iMRI workstation cannot achieve such optimization because of limited hardware resources that are designed only to maximize the performance of the real-time intra-operative image acquisition. The combination of a hardware-accelerated image processing library (XIL, Sun Microsystems, Mountain View, CA) and a compatible graphics board (Creator 3D, Sun Microsystems, Mountain View, CA) processed 2D images much faster in the research workstation than could be done in the iMRI workstation. The accelerated image processing was provided by XIL and includes arithmetic, logic, geometric operations, convolutions, and image statistics. Benchmarking tests showed scaling and convolution functions to have performances of 2517 and 274 images/s (256×256 pixels×16 bits), respectively.

The development platform for the software used Task Command Language/Tool Kit (Tcl/Tk) with a combination of C/C++ languages. Tcl/Tk is an integrated scripting language which not only has most of the capabilities of standard C/C++, but also supports socket level network communication, input-output (I/O) handling, looping, and mathematical manipulation. Tk is the extension of Tcl for graphical use interface construction which manages windows and mouse events. One of the major benefits of Tcl/Tk is rapid prototyping and thus it is simple to develop interactive programs and graphical user interfaces.

Rapid prototyping was an important design decision essential for developing and refining the graphical user interfaces for a physician to use intra-operatively. Both Tcl and Tk have an interface which enables developers to implement custom C/C++ methods such as the two algorithms described below.

2.5.1 Algorithm 1: T1-Weighted FSE Image Subtraction

Intra-operatively acquired difference images provided by the research workstation were generated by subtracting and smoothing consecutive T1-w FSE MR images (Figure 5). The generation of the difference images centered around the subtraction of T1-w FSE images (256×256 pixels×16 bits) consisted of the sequential processes as described:
step (1) image acquisition; access to buffer; data transfer to research workstation;
step (2) FOV definition to enhance the region of interest;
step (3) image subtraction;
step (4) 3×3 low-pass Gaussian filtering for noise reduction;
step (5) user-defined thresholding to remove noise and enhance heated region;
step (6) color coding: red (hot) → Green (warm) → Blue (cool);
step (7) superimpose difference image onto original baseline image (as needed).

The computation time for steps (2)–(7) of the reconstruction process was 27 ms. Step (6) enhanced the visualization of the thermal effect by assigning a pseudo-color based on the values of the difference in the signal intensity from one image to another.

2.5.2 Algorithm 2: Water Proton Chemical Shift

Difference images were generated by subtracting and smoothing consecutive processed chemical shift MR images. The dependence of the water proton chemical shift on temperature is given by:

\[
\Delta \phi = \tan^{-1}\left(\frac{\text{Re}[S(T)] \cdot \text{Im}[S(T)] - \text{Re}[S(T)] \cdot \text{Im}[S(T)\text{b}]}{\text{Re}[S(T)] \cdot \text{Re}[S(T)] + \text{Im}[S(T)] \cdot \text{Im}[S(T)\text{b}]})\right),
\]

where \(\Delta \phi\) is the phase distribution difference between the objective-temperature \(T\) image and the baseline-temperature \(T_b\) image. \(S\) is the complex MR signal; \(\text{Re}\) and \(\text{Im}\) denote the real and imaginary parts, respectively.

The sequential procedure for providing the data intra-operatively for the calculated phase difference images (Figure 5) was nearly the same as shown above for the T1-w FSE images as input. However, here step (3) above was replaced by Eq. (1).

- steps (1)–(2) (same as T1-weighted subtraction);
- step (3) temperature map generation;
- steps (4)–(7) (Same as T1-weighted subtraction).

The total computation time was 221 ms for Eq. (1), which is about ten times longer than a simple subtraction.

3 RESULTS

Patient (JA) was a 76-year-old female with a 2 cm high grade glioma in the left frontal lobe with resulting compression of the left lateral ventricle and midline shift. On the day of LITT, the patient was placed in the iMRI scanner. A clamp was used to fix the position of the head. After establishing a sterile field, a small 2–3 mm skin incision was made and a burr hole drilled through the skull and dura. Under MRI guidance, an MR-compatible sedan-type biopsy needle was placed in the mass and a diagnos-
The current technique used a low-pass Gaussian filtering to highlight the regions of change in the MR signal. However, a simple median filter might be more successful at reducing speckle noise. Alternatively, a more sophisticated edge-preserving smoothing operation could be employed. The penalty for such filtering operations is computation cost, which will thus lengthen the interval between image-feedback updates. Another possible imaging method for navigation of LITT is optical flow analysis.

Although UNIX workstations and network facilities were utilized here to achieve real-time monitoring of LITT, this technology could also be ported to personal computers (PCs). Most of the commercially available PCs and their associated operating systems are capable of performing similar calculations in real-time.

Fig. 6 Images from a case report on 76 year old female, glioma in the posterior left frontal lobe. On the left: T1-w FSE image subtraction. On the right: water proton chemical shift imaging. (a) T1w FSE image (axial, FSE, TR/TE 400/18 ms, slice thickness 5 mm, FOV 220×220 mm, matrix 256×128 pixels, 1 NEX) after the insertion of the guide needle. The white arrow indicates the target lesion and artifact of the guide. (b) T1w FSE image during the ablation. (c) Subtraction image [(b)–(a)] during the ablation. Note high contrast area at the tip of the white arrow around the laser tip. (d) A magnitude image from fast SPGR (TR/TE/flip angle 55/14 ms/20°, slice thickness 4 mm, FOV 320×240 mm, matrix 256×128) before coagulation. (e) Magnitude image from the fast SPGR during the coagulation. (f) Mapping with water proton chemical shift imaging. The intensity level in the black box is equalized for enhanced visibility. “Temperature” distribution is indicated by gray level in the box. The same is also indicated with a black box in (e).

Fig. 7 Same case as in Figure 6: comparison of pre- and post-operative images. Pre-operative image: (a) T1-weighted image (axial, TR/TE 400/16 ms, slice thickness 5 mm, FOV 220×220 mm, matrix 256×128 pixels). A large alteration of intensity is noted in the left frontal and parietal regions with compression of the sylvian fissure. Associated with hypo-intense alteration of the adjacent white matter, a compression of the left lateral ventricle and slight middle shift is seen. (b) Gd-enhanced T1w image (TR/TE 500/16 ms) shows left hemispheric abnormality with a 3 cm area of contrast enhancement. (c) T2-weighted image (TR/TE 3000/95 ms). A significant hyperintense alteration of the white matter surrounds the tumor and extends into the frontal and parietal lobes. Post-operative images 3 days after the LITT. (d) T1-weighted image (TR/TE 400/16) indicates the two laser-induced lesions. Compared to the examination done before LITT, there is a decrease in size of the residual alteration of intensity in the left fronto-shaped area of max. 1.5 cm in diameter. (e) Gd-enhanced T1 (TR/TE 500/16 ms) showed no contrast enhancement at the center of the lesion, although there is an enhancing rim around the tumor site. (f) T2w image (TR/TE 3000/95 ms) has low intensity in the center of the tumor site and high intensity in the rim. The surrounding vasogenic edema has less compression of the left lateral ventricle.
system (e.g., Windows 95) already incorporate networking capabilities. The graphical user interface software also can be ported to a PC environment, as its architecture was carefully chosen to be run across platforms.

5 CONCLUSION

MRI offers a sensitive, noninvasive technology by which interstitial thermal therapies can be monitored and thereby controlled. The application of laser-induced coagulation for tumor therapy has been limited by the ability to monitor the laser-tissue interaction. Slow MRI image acquisition sequences and closed MRI scanners showed the promise of such treatments over the last decade. Now, fast imaging sequences and “open” interventional units deliver on this promise.

As research now looks to optimize the imaging sequences used, there are still important components of such therapies that must be tested and evaluated. A crucial component is the concept of feedback, which is the technological environment by which the feedback is provided. We provided a report on a system design feasible for on-line monitoring of interstitial laser therapy in the brain in a real, dedicated iMRI environment. We described the system’s engineering aspects and shared data from clinical application in a patient.

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