Title
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Permalink
https://escholarship.org/uc/item/09w38944

Journal
Expert review of medical devices, 12(1)

ISSN
1743-4440

Authors
Chen, Clark C
White, Nathan S
Farid, Nikdokht
et al.

Publication Date
2015

DOI
10.1586/17434440.2015.975118

Peer reviewed
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To cite this article: Clark C Chen, Nathan S White, Nikdokht Farid, Peter Kobalka, Lawrence Hansen, Matthew Pearn & Anders M Dale (2015) Pre-operative cellularity mapping and intra-MRI surgery: potential for improving neurosurgical biopsies, Expert Review of Medical Devices, 12:1, 1-5, DOI: 10.1586/17434440.2015.975118

To link to this article: https://doi.org/10.1586/17434440.2015.975118
Pre-operative cellularity mapping and intra-MRI surgery: potential for improving neurosurgical biopsies


Clark C Chen
Author for correspondence:
University of California, San Diego, San Diego, CA, USA
Tel.: +1 858 246 0674
Fax: +1 858 534 8899
clarkchen@ucsd.edu

Nathan S White
University of California, San Diego, San Diego, CA, USA

Nikdokht Farid
University of California, San Diego, San Diego, CA, USA

Peter Kobalka
University of California, San Diego, San Diego, CA, USA

Lawrence Hansen
University of California, San Diego, San Diego, CA, USA

Matthew Pearn
University of California, San Diego, San Diego, CA, USA

Anders M Dale
University of California, San Diego, San Diego, CA, USA

Stereotactic biopsies are frequently performed to secure definitive diagnosis for brain tumor patients. Fundamentally, there are two major difficulties in these endeavors. First, because of intra-tumoral heterogeneity inherent in many forms of brain cancer, biopsies taken from one region may yield a different diagnosis than if another area is biopsied. Second, stereotactic needle biopsies inherently rely on mathematical algorithms for targeting, without real-time visualization of the actual biopsy site. This article describes the novel MRI-based technologies that can potentially afford neurosurgeons the opportunity to address these challenges.

For most solid tumors, definitive diagnosis requires securement of abnormal tissue and careful pathologic examination of this tissue. This tissue diagnosis remains the cornerstone of subsequent therapy [1]. However, there are a number of cancers where intrinsic intra-tumoral histologic heterogeneity render definitive diagnosis challenging. Glioblastoma multiforme, the common form of primary brain cancer, is one such cancer [2].

Glioblastoma is notoriously heterogeneous in terms of the histologic appearances [5]. This heterogeneity exists at a cellular and a regional level, such that biopsy of tissue secured from one region of the tumor may yield a diagnosis of grade III tumor, while specimens secured from another region of the same tumor can render a grade IV diagnosis. In one study, 81 patients afflicted with astrocytomas of differing grades underwent stereotactic biopsy followed by surgical resection (within 30 days), 38% of the initial diagnosis secured through stereotactic needle biopsy were of a grade that differed from diagnosis achieved through surgical resection [9]. In a second study, misdiagnosis occurs in 25% of stereotactic needle biopsies involving lesions <1 cc in volume [10]. These studies demonstrate that misdiagnosis from stereotactic needle biopsies represent a genuine challenge in neurosurgery.
Brain biopsies: challenges

The difficulty of the tissue sampling is confounded by the complexity of the brain, where hundreds of trillions of neuronal connections define those qualities that we consider human [11]. The notion of eloquence is often applied by neurosurgeons to delineate regions of the brain that are amenable to surgical manipulation. In neurosurgery, eloquent cortex is defined by regions where injuries result in obvious neurologic deficits, such as weakness or paralysis. However, there is a large body of literature demonstrating that injuries in regions not considered eloquent, nevertheless, cause deficits that are detectable on sophisticated neurocognitive testing [12,13]. As such, it is our contention that any neurosurgical procedure should be performed with the intent of minimizing injuries to any portion of the cerebrum. As such, it is highly desirable to achieve definitive diagnosis with minimal disruption of the cerebrum.

Another major hurdle in biopsy of the human cerebrum is that it is effectively a ‘blind’ procedure. While sophisticated methods have been developed to triangulate the intended region of the biopsy and to deliver the biopsy needle to this region [10], the neurosurgeon performing the procedure has limited means of visually confirming the location of the biopsy needle. In other words, the accuracy of the biopsy is entirely dependent on precision of the instrument, and the neurosurgeon has little means of validating the accuracy with human aptitudes, such as the surgeon’s experience and intuition. For instance, the stereotactic frame may incur subtle deformation with repeated use. If not properly serviced, utilization of this sub-optimal frame introduces inaccuracies. For this and other well-described factors that influence the accuracy of stereotactic biopsies [14,15], most experienced neurosurgeon will have experienced situations in the operating room (OR) where the frozen specimen analysis is non-diagnostic and the doubts are raised as to the actual location of the biopsy. For the most part, the neurosurgeon makes educated ‘guesses’ in these situations to decide on the subsequent course of action.

The blind nature of brain biopsies additionally impede the surgeon’s ability to react to intraoperative events. For instance, most surgeons would terminate the surgery if they know that the biopsy had triggered significant hemorrhage. However, the way that brain biopsies are currently performed, the surgeons are effectively blind to these events. The surgeon effectively relies on the patient’s neurologic examination as she/he emerges from anesthesia to assess whether adverse events were incurred during the biopsy. If the examination is concerning, the surgeon would then rush the patient to a CT scanner for imaging. The delay between the timing of the actual hemorrhage and the timing of detection on CT can be on the order of hours. Based on the available literature, the risk of biopsy-related hemorrhage ranged between 1 and 9% [16–21].

Improving brain biopsies

Restriction spectrum imaging

The development of diffusion weighted magnetic resonance imaging (DWI) has allowed visualization of microstructural and physiological changes within the brain [22]. The physical principle underlying DWI involves assessing molecular diffusion by the imposition of two radiofrequency pulses that are equal in magnitude and 180 degrees out of phase. The amplitude of the radiofrequency perturbation is characterized by the b value, and most conventional DWI images are acquired using a single b value [23]. Using diffusion imaging, cellularity maps

Figure 1. Restriction spectrum imaging (RSI)-guided selection of biopsy sites. (A) RSI signal imposed onto conventional MR imaging. Red color indicates regions of increased cellularity. Expectedly, the cortex of the cerebrum exhibits increased RSI signal. There is an increased RSI signal on the lateral edge of the tumor mass. (B) Biopsy of a region of tumor with increased RSI signal and another adjacent region without increased RSI signal. (C) Pathologic specimens secured from respective regions. The region of increased RSI signal revealed increased cellularity as well as increased microvascular proliferation. The region without increased RSI signal showed moderate cellularity. Both slides were taken at 20×. Bar = 50 μm. The specimens were stained by H&E.
can be generated using DWI to guide target planning for stereotactic brain biopsies [24].

Restriction spectrum imaging (RSI) is an advanced form of DWI technique that integrates multiple amplitudes of radiofrequency perturbations (hence a spectrum of b values) to assess molecular motion [23]. We have previously shown that RSI afford finer resolution of molecular diffusion relative to conventional DWI, affording assessment of a gradation of diffusional restrictions, ranging from free to partial restriction to absolute restriction [25]. With RSI, we were able to determine regions of high cellularity within the tumor that failed detection by conventional DWI [26]. Incorporation of this information into surgical planning can potentially enhance the surgeon’s ability to select region of disparate cellularity for biopsy (FIGURE 1).

**Intra-MRI biopsies**

With the development of MRI-compatible equipment such as the ClearPoint device [27] and Ad-Tech biopsy needle [28], brain biopsies can now be performed within the MRI. In doing so, the surgeons will have a real-time view of the lesion as it is being biopsied. Adjustments in trajectory can be made in real time to sample the regions of interest.

The ClearPoint device is an integrated system of hardware, software and disposable MRI compatible instruments that afford surgeons a real-time view of the surgical lesion and the biopsy needle in real time as the biopsy is being performed. To the best of our knowledge, it is the only commercial device that allows for real-time MRI-guided neuronavigation. The patients undergoing a ClearPoint procedure are placed under general anesthesia. A set of MRI images were then taken and used for planning surgical trajectory. Based on this trajectory, an incision is made followed by a dime-sized burr hole through the skull. A tripod device (termed SmartFrame) is mounted over the incision (FIGURE 2). This frame is synchronized to the hardware and software such that the trajectory of the biopsy needle inserted through the center of the tripod can be calculated. Based on this calculation, the needle is advanced slowly to the intended target. MRIs are performed during this advancement as well as during the actual biopsy to visualize the trajectory in real time. Because actual views of the process are
available in real time, the surgeon can use his/her judgment to adjust to any inaccuracies related to the surgical equipment and react to any intraoperative events encountered.

Importantly, performing the biopsy in the MRI allows the neurosurgeon to more quickly react to the situation of an enlarging hematoma. For deep-seated tumors, the management strategy for an expanding hematoma involves termination of biopsy, correction of aberrant coagulation parameters and blood-pressure control, with surgical evacuation in the case of hematoma exerting significant mass effect. To determine whether hemorrhage has occurred and the size of the hematoma, the patients are typically emergently transported from the OR (where conventional biopsies are performed) to the imaging suite. If the hematoma size is significant, the patient is then brought back to the OR. In the case of MRI-guided biopsies, the patient is already under surveillance by MRI and an abbreviated T2 sequence would determine whether the hematoma is of a size that requires evacuation, thereby bypassing the trip from the OR to the CT. Thus, the proper course of action can be more quickly determined for a patient biopsied in the MRI relative to the patient who was biopsied in the OR.

A link to a video describing the integration of pre-operative RSI planning and intra-MRI biopsy can be found in Reference [29]. The video describes a case where regions of differing RSI signals were biopsied with the MRI. The localization of the regions of biopsies was visually confirmed in real time. The biopsy derived from a region of lower cellularity (or low RSI signal) yielded the diagnosis of anaplastic astrocytoma. In contrast, the biopsy derived from a region of high cellularity (or high RSI signal) yielded the diagnosis of glioblastoma (Figure 1). The case illustrates the potential of RSI imaging and real-time MRI in improving the accuracy of stereotactic needle biopsies.

**Conclusion**

Technological advances have now conferred neurosurgeons with the ability to pre-operatively define the regional heterogeneity of brain tumors as well as real-time visualization of biopsy as it is performed. Adaptation of these technologies can potentially improve the safety and accuracy of brain biopsies. However, assessment of efficacy, cost–benefit analysis and clinical experience from larger cohorts are needed before clinical adaptation of these technologies.

**Financial & competing interests disclosure**

The authors have no relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript. This includes employment, consultancies, honoraria, stock ownership or options, expert testimony, grants or patents received or pending, or royalties.

No writing assistance was utilized in the production of this manuscript.

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