Presurgical Brain Mapping of the Ventral Somatomotor Network in Patients with Brain Tumors Using Resting-State fMRI


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ABSTRACT

BACKGROUND AND PURPOSE: Resting-state fMRI readily identifies the dorsal but less consistently the ventral somatomotor network. Our aim was to assess the relative utility of resting-state fMRI in the identification of the ventral somatomotor network via comparison with task-based fMRI in patients with brain tumor.

MATERIALS AND METHODS: We identified 26 surgically naive patients referred for presurgical fMRI brain mapping who had undergone both satisfactory ventral motor activation tasks and resting-state fMRI. Following standard preprocessing for task-based fMRI and resting-state fMRI, general linear model analysis of the ventral motor tasks and independent component analysis of resting-state fMRI were performed with the number of components set to 20, 30, 40, and 50. Visual overlap of task-based fMRI and resting-state fMRI at different component levels was assessed and categorized as full match, partial match, or no match. Rest-versus-task-fMRI concordance was calculated with Dice coefficients across varying fMRI thresholds before and after noise removal. Multithresholded Dice coefficient volume under the surface was calculated.

RESULTS: The ventral somatomotor network was identified in 81% of patients. At the subject level, better matches between resting-state fMRI and task-based fMRI were seen with an increasing order of components (53% of cases for 20 components versus 73% for 50 components). Noise-removed group-mean volume under the surface improved as component numbers increased from 20 to 50, though ANOVA demonstrated no statistically significant difference among the 4 groups.

CONCLUSIONS: In most patients, the ventral somatomotor network can be identified with an increase in the probability of a better match at a higher component number. There is variable concordance of the ventral somatomotor network at the single-subject level between resting-state and task-based fMRI.

ABBREVIATIONS: BOLD = blood oxygen level–dependent; ICA = independent component analysis; rs-fMRI = resting-state fMRI; tb-fMRI = task-based fMRI; VSMN = ventral somatomotor network; VUS = volume under the surface

Functional MRI is widely used as a noninvasive tool for presurgical localization of the eloquent cortex, typically involving somatomotor and language mapping. Mapping of these eloquent brain areas with fMRI correlates well with invasive methods such as intraoperative electrocortical stimulation and can result in reduced surgical time, increased extent of resection, and decreased craniotomy size. In addition, postoperative morbidity correlates with the distance of the resection margin from fMRI-identified eloquent cortex. In current clinical practice, changes in blood oxygen level–dependent (BOLD) signal are measured across time as the patient performs a specific task (ie, task-based fMRI [tb-fMRI]). These BOLD signal changes reflect characteristic hemodynamic responses to neural activity. However, tb-fMRI has several limitations. Primarily, accurate localization of function is dependent on the patient’s cooperation and ability to...
adequately perform the task, which can be an important limitation in those with physical or mental debilitation or those who cannot understand the instructions such as children. The patient must be awake and cooperative during the task; therefore, sedation cannot be used; this feature is a limitation in the pediatric population. Furthermore, due to the inherently low signal-to-noise ratio of this technique, long and repeated acquisitions are often required to ensure adequate data sampling for analysis.5,6

Resting state fMRI (rs-fMRI) is a promising method of assessing brain function that can overcome some of the limitations of tb-fMRI. In rs-fMRI, spontaneous fluctuations in BOLD signal are measured across time while no specific task is performed. Temporal correlations of these spontaneous fluctuations can be organized into spatially distinct intrinsic networks.5 These networks are now described as resting-state networks, many of which have topography similar to that of networks engaged in sensory, motor, and cognitive tasks.7 These intrinsic resting-state networks persist, though somewhat modified, in states of decreased awareness such as sleep8 or sedation.9 Rs-fMRI has thus gained interest as a potential viable alternative to tb-fMRI, especially in pediatric or cognitively/neurologically impaired patients. Multiple studies have investigated the potential role of rs-fMRI in presurgical mapping of somatomotor5,10-12 and language2,6,13 networks. In a limited number of subjects, moderate overlap between rs-fMRI and tb-fMRI was found for mapping of the motor cortex.5,11,12 Furthermore, there is good qualitative concordance between intraoperative cortical stimulation and rs-fMRI in the localization of the eloquent motor cortex.10 With quantitative analysis, rs-fMRI and tb-fMRI perform comparably, but the shortest distance to stimulation points is observed for tb-fMRI.11 Rs-fMRI has been shown to identify a larger pattern of the motor network compared with tb-fMRI.11,14 In addition, localization by rs-fMRI and tb-fMRI may include different parts of the somatomotor network.11,15 While the motor system at large has been the target of most prior investigations, distinct subnetworks of the motor network exist. We narrowed our focus to the ventral motor area, which largely reflects oral somatomotor function. Three recent studies evaluated the concordance of rs-fMRI and tb-fMRI in the localization of the face representation area of the primary motor cortex in addition to limb motor areas in a limited number of patients with a variety of brain lesions in different locations.11,12,16

To our knowledge, no study has specifically investigated the concordance between tb-fMRI activation and the rs-fMRI-derived ventral somatomotor network (VSMN) maps in a large cohort of patients with lesions close to or involving the VSMN. We hypothesized that the ventral motor network can be identified in patients with brain tumors and that there is good concordance with tb-fMRI with a tongue motor paradigm. In addition, we hypothesized that a higher number of components in independent component analysis (ICA) may yield a better concordance between rs-fMRI and tb-fMRI.

MATERIALS AND METHODS

Study Subjects
The institutional review board approved this retrospective study. Searching the Radiology Information Systems, we identified patients who underwent fMRI for presurgical brain mapping between January 1, 2009, and July 31, 2014. Fifty-eight patients underwent rs-fMRI in addition to tb-fMRI during the same imaging session. Seventeen patients had a prior history of a brain operation (including biopsy) and were excluded. One patient had imaging features characteristic of an arteriovenous malformation and was excluded. At the Johns Hopkins Hospital, tb-fMRI for presurgical mapping is tailored for each patient, predominantly based on the location of their tumor and the relation of eloquent regions along the expected surgical trajectory and in the immediate vicinity of the lesion. Among the remaining 40 patients, 26 with brain tumors involving or in close proximity to the ventral somatomotor network who had undergone both tb-fMRI for localization of the VSMN and rs-fMRI were included for final analysis (age range, 21–69 years; mean age, 43.6 years; 15 men and 11 women).

Brain Tumor Characterization
Tumor location, volume (in cubic millimeters), pathology, and World Health Organization histologic grade (when available) were recorded by a subspecialty board-certified neuroradiologist for each patient. Lesion volume was measured by manually drawing the ROI on FLAIR images in the Medical Image Processing, Analysis, and Visualization application software (MIPAV; National Institutes of Health, Bethesda, Maryland; http:// mipav.cit.nih.gov). For high-grade brain tumors, we assessed the entire region of T2 signal abnormality, which represents a combination of infiltrative neoplasm and vasogenic edema.

MR Imaging
A 3T Tim Trio system (Siemens, Erlangen, Germany) with a 12-channel head matrix coil was used. Structural images included a 3D T1 sequence (TR = 2300 ms, TI = 900 ms, TE = 3.5 ms, flip angle = 9°, FOV = 24 cm, acquisition matrix = 256 × 256 × 176, section thickness = 1 mm) and a 2D T2 FLAIR sequence (TR = 9310 ms, TI = 2500 ms, TE = 116 ms, flip angle = 141°, FOV = 24 cm, acquisition matrix = 320 × 240 × 50, section thickness = 3 mm). Functional T2*-weighted BOLD images for both tb-fMRI and rs-fMRI were acquired by using 2D gradient-echo echo-planar imaging (TR = 2000 ms, TE = 30 ms, flip angle = 90°, FOV = 24 cm, acquisition matrix = 64 × 64 × 33, section thickness = 4 mm, section gap = 1 mm, interleaved acquisition). Instructions for rs-fMRI were the following: try to keep still, keep your eyes closed, and do not fall asleep. For rs-fMRI, 180 volumes were acquired (6 minutes).

Tongue Motor Task
The face representation area of the primary motor cortex was mapped by using the vertical tongue movement task as a robust and spatially extensive representative of the ventral/facial motor network.17,18 The duration of the task is 3 minutes and consists of 3 cycles of 30-second blocks of rest alternating with 30-second blocks of repetitive vertical tongue movement. For each patient, a board-certified neuroradiologist with experience in fMRI provided instructions and practice sessions on the tongue motor task outside the scanner. The quality of the tb-fMRI maps was monitored in real-time by the neuroradiologist monitoring the session. Per protocol, if real-time maps dem-
onstrated suboptimal activation or excessive noise, the task was repeated. Tasks were presented with the environment implemented in the Prism Acquire software (Prism Clinical Imaging, Elm Grove, Wisconsin). Only runs that met quantitative quality control criteria (≤2-mm net head displacement along any axis) were considered.

Image Processing
fMRI data were processed by using Statistical Parametric Mapping, Version 8 (SPM8 software; http://www.fil.ion.ucl.ac.uk/spm/software/spm12) and custom Matlab (MathWorks, Natick Massachusetts) scripts. Processing of tb-fMRI included section-timing correction, motion correction, normalization to a Montreal Neurological Institute 152 template, and spatial smoothing included a 6-mm full width at half maximum Gaussian kernel.

For processing the rs-fMRI data, section-timing correction and motion correction were performed. The motion-correction step included registration of tb-fMRI and rs-fMRI to each other. The ArtRepair toolbox (http://cibsr.stanford.edu/tools/human-brain-project/artrepair-software.html)19 was used to detect volumes with large shifts in global average signal intensity, which include contributions from scan-to-scan motion. The outlier volumes and additional volumes recommended for deweighting in ArtRepair were tagged for subsequent removal from analysis (ie, scrubbing). The rs-fMRI data were then linearly detrended. Rs-fMRI and T1-weighted images were coregistered and normalized to perform physiologic nuisance regression of rs-fMRI with the component-based noise-correction method.20 The same transformation matrix was used for normalization between tb-fMRI and rs-fMRI to ensure that spatial comparison between these 2 was valid for each subject. Bandpass filtering from 0.01 to 0.1 Hz and smoothing were performed with a 6-mm full width at half maximum Gaussian kernel. At the end, previously tagged images by ArtRepair were scrubbed.

Statistical Analysis
Tb-fMRI Analysis. We used a general linear model analysis for tb-fMRI implemented in SPM8 with the canonical hemodynamic response function convolved with the boxcar function for each task with standard parameters previously described.6 High-pass filtering was performed at the default setting to remove drift. The hemodynamic response function-convolved task vectors were input into a design matrix, and a contrast was created to target the associated parameter comparing the hemodynamic response function-weighted time engaged in the task with rest. The SPM t-contrast maps were generated without clustering because these activation maps were subsequently thresholded across multiple levels for comparison as previously described.6 Activation maps were reviewed to ensure that ventral motor activation was present.

Rs-fMRI Analysis. Rs-fMRI was analyzed with the Group ICA of the fMRI Toolbox Software (GIFT; http://mialab.mrn.org/software/gift/). Independent component analysis was performed separately for each subject by using the InfoMax algorithm with ICASSO (http://research.ics.aalto.fi/ica/icasso/) set at 5 repeats,21 including selection of the “best run” to ensure consistent estimates. ICA maps were generated for 20, 30, 40, and 50 components, designated here as ICA20, ICA30, ICA40, and ICA50, respectively. Following scrubbing, 1 subject had only slightly >50 volumes left; therefore, the maximum number of components was limited to 50. For each ICA group, the component that best represented the VSMN based on overlap with tb-fMRI was selected visually (Fig 1). Any component whose spatial map was specific and limited to the ventral perirolandic cortices corresponding to the tb-fMRI activation maps was considered a “full match.” If the component included additional networks outside these regions, it was considered a “partial match,” indicating that the VSMN was present but not separated from other networks. If no VSMN was identified, a “no match” designation was assigned to that ICA group for that subject. A mixed-effects logistic regression analysis was performed to see whether there was a significant increase in the probability of getting a full match as a function of the number of ICA components.

Comparison of tb-fMRI and rs-fMRI. The level of tb-fMRI T-map thresholding may affect the degree of rs-tb fMRI concordance. At low thresholds, there will be artificially high concordance due to the introduction of a higher percentage of voxels, many of which may not represent true activation but rather statistical noise that
exceeds this threshold. Although there are strategies to determine the level of optimum fMRI thresholding, there is no consensus for reproducible results at a subject level. To minimize this issue, we used a multithresholding technique to compare concordance across a wide range of thresholds as previously described (Fig 2). We calculated Dice coefficients as a quantitative measure of overlap across each threshold to generate a matrix of rs-versus-tb overlap (left map, Fig 3). Dice coefficients vary between 0 and 1 and give an objective evaluation of similarity or concordance between 2 sets of data. At very low thresholds, high Dice coefficients result from overlap of random noise as can be seen in Fig 2. We used a previously described noise-removal method briefly described here.

For calculation of noise, an ICA component representing the anterior ventricular signal was selected for each subject. After removing negative values, we normalized image maps in value from 0 to 1, and we subsequently used multiple thresholds of this normalized map to calculate Dice coefficients between ventricular 

“noise” and task maps (middle map, Fig 3). The resultant noise-versus-task Dice map was subtracted from the resting-versus-task Dice map (right map, Fig 3). Noise-corrected resting state-versus-task-based Dice map volume under the surface (VUS) was calculated across different ICA orders. The VUS measurement collapses this multithresholded dice map into a single variable that can be used as a metric; the VUS is equivalent to the area under the curve in a 2D graph (such as a receiver operating characteristic curve); however, because the multithresholded dice map has 3 dimensions (the rs-fMRI threshold, the task-threshold, and value), a volume under the surface is computed. One-way ANOVA was performed to determine significant differences in VUS across the 4 ICA orders. Maximum Dice coefficient values at the group level and subject level were calculated.

RESULTS
Tumors
The On-line Table summarizes the patients’ demographic data, location and volume of lesions, pathology, World Health Organization grade (if applicable), and the distance from the edge of the lesion to the edge of the ventral somatomotor activation cluster based on the clinical task-fMRI maps. Brain lesions were mainly centered in the left cerebral hemisphere (19 patients), and 4 lesions were primarily located in the right cerebral hemisphere. Tumor size ranged from 0.84 to 159.05 cm³ (mean, 41.82 cm³).

FMRI Comparison
Twenty-six patients met the inclusion criteria and were included for analysis. In 21 patients (81%), rs-fMRI successfully identified a VSMN (Fig 4). Among these, 2 patients were considered a partial match because the VSMN was not distinct from the dorsal somatomotor network. There was an increase in the number of full matches with an increase in the ICA order (14 patients at ICA 20 and 19 patients at ICA 50). In 14 patients, a separate VSMN was identified at all ICA groups. Mixed-effect logistic regression demonstrated a significant increase in the probability of getting a full match as a function of the number of components (< .00001). This probability is shown in Table 1.

In 5 patients, rs-fMRI failed to identify a VSMN at any of the ICA levels. In 3 of these 5 patients, only the dorsal somatomotor

FIG 2. Rs-fMRI ICA maps (yellow) and Tb-fMRI T-maps (red) were thresholded, and 100 threshold maps were generated for each in a single subject. At a low threshold, there is artificially higher overlap (orange) between the maps due to noise. At a very high threshold, the overlap is smaller.

FIG 3. The Dice coefficient matrix at different thresholds at the subject level between the rs-fMRI (x-axis) at ICA 20 and tb-fMRI (y-axis). An artificially high Dice coefficient is seen in the top left corner of the left map due to overlap of noise. A noise matrix was generated (middle map) and was subtracted (right map). Noise-removed Dice coefficient maps were generated for all the subjects across 4 different ICA orders.
network was identified. In the remaining 2 patients, no somatomotor network was identified.

Group mean Dice maps for each ICA order are shown in Fig 5. Group mean Dice VUS overall increased with the ICA order (Fig 6); however, 1-way ANOVA demonstrated no significant differences among the 4 ICA groups (P value = .4). The range of Dice coefficient values at the subject level is shown in Table 2. Negative Dice values in these noise-subtracted maps may occur when the concordance between the noise maps and tb-fMRI at low thresholds is greater than the concordance between rs-fMRI and tb-fMRI due to the randomness of noise at low thresholds.

**DISCUSSION**

A growing number of studies have explored the feasibility of rs-fMRI as a substitute or complement to tb-fMRI for presurgical mapping of the eloquent somatomotor cortex in patients with brain tumor. Previous studies have demonstrated the reliability of rs-fMRI for localization of the hand motor area in comparison with tb-fMRI in a limited number of patients, with seed-based or ICA analysis. Concordance of rs-fMRI with intraoperative cortical stimulation was assessed in a few patients. In addition, investigators have assessed the entire somatomotor cortex by using hand, foot, and face paradigms, with evaluation of the face motor area in a small subset of patients. A novel data-driven algorithm for mapping the functional cortex in preoperative planning of motor and language networks in a small group of patients with tumor and epilepsy has been reported. The existing data predominantly evaluate the usefulness of rs-fMRI in localization of the hand representation area of the somatomotor cortex, but there has been very limited evaluation of the ventral somatomotor cortex to date. In addition, there is wide variation in analysis methods, and most studies have only qualitatively compared tb-fMRI and rs-fMRI. To our knowledge, our study represents a relatively large cohort of patients with brain tumors involving or in close spatial proximity to the VSMN. In addition, we evaluated the overlap quantitatively by using multithresholding at different ICA levels.

![FIG 4. fMRI comparison among the patients. The number of patients with full, partial, and no match between tb- and rs-fMRI at different ICA levels is demonstrated.](image1)

![FIG 5. Group mean Dice coefficient maps across all the subjects at each ICA order calculated from subject-level noise-removed Dice coefficient maps. The x-axis depicts the rs-fMRI threshold levels, and the y-axis depicts the tb-fMRI threshold levels with the color bar demonstrating the Dice coefficient value.](image2)

![FIG 6. Violin plots of Dice VUS across ICA orders. The mean Dice VUS for each ICA order is denoted by white diamonds. As the ICA order increases, there are larger numbers of subjects with higher Dice VUS values as reflected in the greater width of the violin plots corresponding to kernel densities.](image3)

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<th>Table 1: Probability of getting a full match between rs-fMRI and tb-fMRI maps as a function of number of ICA components</th>
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<td>ICA 20</td>
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<td>Probability</td>
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<th>Table 2: Dice coefficient values range, median, and mean at the subject level across different ICA orders</th>
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<td>ICA 20</td>
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place placement of seeds may be limited due to anatomic gyral distortion or compression. Therefore, an unbiased method such as ICA may be preferable. In addition, ICA remains an attractive choice due to availability of easy-to-use software such as the MELODIC tool in FSL (http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/MELODIC) or the GIFT toolbox.

Several limitations in this study should be addressed. rs-fMRI was performed after tb-fMRI in this clinical cohort of patients to ensure that the patients tolerated the lengthy scan and performed well on their tb-fMRI, which was critical for clinical presurgical mapping. Thus, there may have been an inadvertent task effect on the observed functional connectivity.29 Another limitation is the inability to ensure that patients did not fall asleep during rs-fMRI. We instructed the patients to stay awake during the rs-fMRI and confirmed that they stayed awake during a postscan interview; however, the accuracy of their statements could not be verified by physiologic measures indicating sleep during the acquisition. While this inability poses a potential limitation because changes in functional connectivity have been reported in sleep or altered consciousness,30 functional connectivity in the somatomotor network has been shown to be preserved during different states of arousal.6 Nevertheless, the effect of sleep or altered arousal on rs-fMRI concordance with tb-fMRI could be further investigated. In addition, we used tb-fMRI T-maps to find the best candidate ICA map to represent the VSMN. However, brain tumors may cause alteration of somatomotor network organization, and brain tumor–induced neurovascular uncoupling may further compromise our ability to accurately detect the VSMN. Therefore, using only rs-fMRI data to identify the best somatomotor map may be challenging with brain tumors. The use of a data-driven neural network algorithm to identify the eloquent cortex in 7 patients with brain tumor with distorted anatomy has shown some promise.25

CONCLUSIONS

We demonstrate variable concordance of rs- and tb-fMRI at the single-subject level for detection of the VSMN in patients with brain tumor. We demonstrate improved reliability of rs-fMRI VSMN maps with higher ICA orders. Failure of rs-fMRI to identify the VSMN in about one-fifth of patients in our study limits the ability of rs-fMRI to completely substitute for tb-fMRI for presurgical assessment, though it may be considered as a supplement for tb-fMRI, for example, when task activation is suboptimal. Further improvements in technique, processing, and analysis methods may enhance the potential of using rs-fMRI as a substitute or complement to tb-fMRI in mapping of the VSMN.

FIG7. Sample subject (patient 23 in the On-line Table) demonstrating an expansile mass lesion centered in the postcentral gyrus (arrows). Signal abnormality is extended to the subcortical white matter of the precentral gyrus. Red denotes tongue motor task activation, green denotes the VSMN network identified from rs-fMRI (ICA 50), and yellow denotes areas of overlap between tb- and rs-fMRI.
REFERENCES


