Neurophysiological evidence of preserved connectivity in tuber tissue

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Resective surgery is the only curative option for medically refractory epilepsy in TSC, and >60% of TSC patients with a well localized seizure focus become seizure-free after tuberectomy [3]. Such clinical success motivates exploration into whether and to what extent tubers may have functional connectivity within the brain and spinal cord. Cortical tubers evident on conventional MRI have long been deemed inert, static lesions that can be safely resected [3]. However, recent neurophysiology, neuropathology, and diffusion imaging demonstrate absence of clear tuber-cortex boundaries, with cortical elements retained within the tuber and extensive tuber-like pathology beyond the tuber [4]. Relevant to the present report, tubers contain maloriented pyramidal cells and dysplastic neurons that suggest a tuber-to-cortex connectivity [5]. Further, tubers, like focal cortical dysplasias, can be in the proximity of eloquent cortex, leading to investigations into whether tubers can contain function [3].

Notably, it is unknown if structural connectivity or neurophysiologic function can be preserved in such imaging-evident lesions, where neuropathology is intermixed with normal cortex. The lack of clear MRI-defined resection margins, either on tuber interface with cortex or with normal-appearing white matter (NAWM), poses a challenge as long-term success of epilepsy surgery in TSC is directly determined by the total extent of the resection [6]. Further, as more lesions are detected by increasingly high resolution MRI, the
Neuropsychological testing revealed a 4 year delay behind expected age. He did not have the capacity to write letters or recognize numbers. While the patient was able to use a crayon, he did not have the fine motor skills to draw small objects. Additionally, the patient showed problems with the coordination of complex motor movements. Furthermore, the patient tested positive for a Tsc2 gene mutation.

The patient underwent functional mapping to assess the cortical representation of the left hand. DTI tractography was performed. The results showed evidence of functional connectivity within tuber tissue.

The patient's MRI DICOM file was uploaded into the Nexstim NBS 4.3 software (Nexstim™, Helsinki, Finland) to create a three-dimensional reconstruction of the patient's cortical surface (Fig. 1). Stimulation was performed with a Nexstim™ unit and a figure-of-eight coil operated in the figure-of-eight mode. Motor threshold was operationally defined as the minimum machine output necessary to elicit a response from the APB, deltoid, and TA by surface EMG.

Verbal and written consent were obtained from the patient's parents prior to stimulation. Electromyography (EMG) was recorded through surface electrodes placed in 6 locations on both the right and left sides of the body over the bilateral abductor pollicis brevis (APB), deltoid, and tibialis anterior (TA) muscle groups. An additional ground electrode was placed on the underside of the right forearm. With single pulse nTMS, stimuli were applied at scalp sites overlaying the motor cortex in the right and left hemispheres, while MEPs were recorded bilaterally from the APB, deltoid, and TA by surface EMG.

The patient's MRI showed subependymal nodules, subcortical tubers, and the largest of which was in the left central area. 3T MRI with 35 direction DTI showed charges in left central and frontocentral areas, and seizures originating in the left central area. 3T MRI with 35 direction DTI revealed subependymal nodules, radial migration lines, and multiple subcortical tubers, in part mineralized, and the largest of which was in the left frontal-parietal area. 2-fluoro-deoxy glucose positron-emission tomography (2FDG PET) showed multiple bilateral foci of decreased cortical isotope uptake corresponding to the cortical tubers seen on MRI. Ictal SPECT revealed relative increased perfusion focally in the left frontal lobe when compared to the interictal study. Thus, the active seizure focus was localized to the left frontotoparietal tuber complex, and the patient underwent functional mapping to assess the cortical representation of both left and right hemispheric motor control regions.

2. Methods

2.1. Clinical summary

A 10-year-old, left-handed boy with TSC and intractable epilepsy was referred to Boston Children's Hospital Comprehensive Epilepsy Program for consideration of epilepsy surgery. At birth, he was found to have a cardiac murmur, and echocardiogram showed cardiac rhabdomyomas. A subsequent MRI of the brain revealed classic stigmata of TSC, including subcortical tubers and subependymal nodules. He tested positive for a Tsc2 gene mutation.

Developmentally, the patient's vocabulary was limited to one word by 2 years of age. He began walking at 18 months with difficulties in coordination of complex motor movements. Additionally, the patient struggled with fine motor skills such as effectively using utensils. At the time of presentation he was able to speak in short sentences and was capable of expressing desire. While the patient was able to use a pencil to scribble on paper, he did not have the capacity to write letters. Neuropsychological testing revealed a 4 year delay behind expected cognitive age. Physical examination was remarkable for hypomelanotic macules on the right lower leg, left thigh and left chest, a shagreen patch on the right medial scalpula, and facial angiofibromas. On neurological exam he appeared left-handed, but was ambidextrous for some tasks. He had a low central tone but no lateralizing motor findings.

His seizure disorder first presented with infantile spasms, which were successfully controlled with vigabatrin. He then developed focal dyscognitive seizures refractory to multiple anti-epileptic medications, with variable seizure semiology overt time. During evaluation in the epilepsy monitoring unit, seizures began with quivering of the chin, and parent-report stated the patient appeared very frightened. Occasionally bilateral oromotor and hand automatisms were seen during which the patient continued to speak with his mother. EEG revealed interictal discharges in left central and frontocentral areas, and seizures originating in the left central area. 3T MRI with 35 direction DTI revealed subependymal nodules, radial migration lines, and multiple subcortical tubers, in part mineralized, and the largest of which was in the left frontal-parietal area. 2-fluoro-deoxy glucose positron-emission tomography (2FDG PET) showed multiple bilateral foci of decreased cortical isotope uptake corresponding to the cortical tubers seen on MRI. Ictal SPECT revealed relative increased perfusion focally in the left frontal lobe when compared to the interictal study. Thus, the active seizure focus was localized to the left frontotoparietal tuber complex, and the patient underwent functional mapping to assess the cortical representation of both left and right hemispheric motor control regions.

2.2. nTMS

Verbal and written consent were obtained from the patient's parents prior to stimulation. Electromyography (EMG) was recorded through surface electrodes placed in 6 locations on both the right and left sides of the body over the bilateral abductor pollicis brevis (APB), deltoid, and tibialis anterior (TA) muscle groups. An additional ground electrode was placed on the underside of the right forearm. With single pulse nTMS, stimuli were applied at scalp sites overlaying the motor cortex in the right and left hemispheres, while MEPs were recorded bilaterally from the APB, deltoid, and TA by surface EMG.

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2.2.1. Data acquisition

The precentral gyrus (M1) and surrounding supplementary motor areas were stimulated to create a map depicting the areas of the cortex...
responsible for eliciting MEPs for a given muscle. The patient had a very high resting APB and deltoid motor threshold (rMT), approximating 100% machine output (MO). Accordingly, upper limb motor mapping was conducted at 100% MO, as opposed to standard laboratory protocol of stimulating at 120% rMT. TA MEPs could not be elicited at rest, and therefore lower limb motor mapping was conducted with the TA muscles activated by foot dorsiflexion, also at 100% MO – this maneuver was sufficient to enable reliable TA MEPs.

2.2.2. Data processing
Results were reviewed by visual inspection for MEP quality to ensure each individual MEP morphology, latency and distinction from background noise were adequate to designate each signal as a valid motor response to stimulation. To visualize the motor mapping results a voltage-graded color scale was generated per recorded MEP such that lower voltages were shown in red and the highest voltages in white. The color scale was generated separately for each muscle group for minimum and maximum values.

2.3. DTI
Imaging was performed for clinical indication on a Siemens Trio 3T MRI system. Sedation was used to prevent significant motion. MR image analysis was done using the CRKit (http://crl.med.harvard.edu) and visualization was performed using the Fibernavigator [9] (https://github.com/scilus/fibernavigator).

2.3.1. Data acquisition
Imaging acquisition parameters are summarized in Table 1. Compensation for residual distortion and patient motion was achieved by rigid registration of the diffusion images to the T1-weighted MPRAGE scan, with appropriate reorientation of the gradient directions [10]. Tensors were estimated using robust least squares [11].

2.3.2. Data processing
The intracranial cavity was segmented into white matter (WM), gray matter (GM), cortex and cerebrospinal fluid using a previously validated segmentation algorithm, based on the individual T1-weighted image [12–14]. Multi-fiber model estimation was done using Distribution of Anisotropic Microstructural eNvironments in DCI (DIAMOND) [15], resulting in up to 3 main peaks per voxel. With this model, we were able to achieve a sharp representation of the fascicles at each voxel, while maintaining sensitivity to the micro-structural integrity of each fascicle.

2.3.3. Tractography
Whole-brain streamline tractography was performed using real-time multi-peak tractography [9], allowing for interactive visualization of streamlines during the tracking process. Our tractography algorithm employs information from the previous step during tracking, based on an earlier DTI-based approach called tensor deflection [16–18], known to ameliorate neurosurgical tracking [19]. Tractography parameters were set as follows: step size: 1 mm, θmax: 45°, streamline min/max length: 60/200 mm, 1 seed per voxel. WM and GM segmentations were used as tracking masks [20], as they provide an improvement

<table>
<thead>
<tr>
<th>Sequence</th>
<th>Voxel size (mm)</th>
<th>FOV (cm)</th>
<th>TE (ms)</th>
<th>TR (ms)</th>
<th>flip angle</th>
<th>acquisition matrix</th>
<th>other</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1-weighted MPRAGE</td>
<td>1 × 1 × 1</td>
<td>19.2–25.6</td>
<td>1.66–3.39</td>
<td>1113–2530</td>
<td>7–9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>T2-FLAIR Turbo spin echo sequence</td>
<td>0.9 × 0.9 × 0.9</td>
<td>19–26</td>
<td>390–400</td>
<td>5000</td>
<td>20</td>
<td>256 × 256</td>
<td>n(excitations) = 1, ETL 141</td>
</tr>
<tr>
<td>Diffusion Single shot SE</td>
<td>1.72 × 1.72 × 1.72</td>
<td>22</td>
<td>88</td>
<td>10</td>
<td>90</td>
<td>128 × 128</td>
<td>30 b = 1000 s/mm2, 5b = 0 s/mm2 images*</td>
</tr>
</tbody>
</table>

* Twice refocused gradients to minimize Eddy currents, iPAT 2, in-plane GRAPPA, modified as necessary to facilitate completion of the scan.
compared to fractional anisotropy (FA)-based tracking methods where streamline propagation is often prematurely halted in crossing regions [21–23]. Streamlines were allowed to propagate freely inside the WM, and up to a maximum of 5 consecutive steps once they reached the GM. Finally, 2D streamlines maps were generated using track-density imaging (TDI) [24].

3. Results

3.1. Motor map results

Well-formed right MEPs were elicited with stimulation over M1, as expected, in the left, contralesional hemisphere. However, in the right hemisphere, robust MEPs from the left TA and deltoid were seen with stimulation almost exclusively over a tuber located in the central sulcus, between pre- and postcentral gyrus. Composite maps for our target muscles are shown in Fig. 2. In contrast to the deltoid and TA muscles, the left APB was mapped to the expected hand region of the right precentral gyrus, outside of the tuber tissue.

3.2. DTI results

Streamlines from the DTI tractography of the corticospinal tract (CST) are preserved in the tuber tissue, as shown in Fig. 3. FA values inside the tuber lesion were approximately 0.35, more than sufficient for streamline fascicle propagation using an FA-based approach with a stop criterion. Visual inspection of the diffusion profiles orientation within the tuber showed coherent organization with respect to the peaks forming the CST.

4. Discussion

Our presurgical nTMS motor mapping results indicate that corticospinal (CS) connectivity may be preserved within the volume of some tubers. Specifically, for a cortical tuber located within the central sulcus, (1) MEPs were elicited directly with stimulation of this area, and (2) these MEPs were of similar amplitude and morphology as MEPs elicited in non-tuber cortex. The functional, physiological evidence of connectivity was further corroborated by DTI tractography showing intact CS connectivity within the tuber tissue.

4.1. Neuropathology and function are intermixed in TSC

The success of resective surgery in TSC patients depends upon the accurate identification of the epileptogenic region [6]. Resection of most epileptogenic tubers, perhaps due to a high degree of intra-tuber pathology, appears to be safe with respect to likelihood of acquired neurologic deficit after surgery [3]. Notably, by MRI protocols similar to those in the present report, large tubers appeared to be circumnavigated on DTI tractography [25]. However, our case suggests that CS connectivity may remain intact in some cortical tubers. Further, we show that nTMS in TSC, as in other disorders where functional mapping is required before neurosurgery, can provide insight into local function of targeted tissue, and may also contribute to the prognosis for post-surgical neurologic deficit. (Note that the tuber with preserved CS connectivity was not the surgical target in this case.)

4.2. Incomplete reallocation of function

Early developmental lesions including stroke and malformations of cortical development may allow for transfer of function to adjacent gyri or to more remote regions [8]. Yet, such functional “remapping” was not identified in our case. The lesion burden of the tuber located in the right central sulcus was sufficient to be evident on conventional MRI, but insufficient to disrupt or warrant transfer of motor function. There is no existing literature discussing the possibility of intact motor function within a tuber. Our case begins to address this gap in knowledge. We thus hypothesize that tubers with a relatively small amount of pathology may contain function. The observed physiologic and imaging phenomena, the presence of robust MEPs from nTMS stimulation and preserved CS tracts identified by DTI tractography, within the tuber tissue in our patient provide functional and structural support of the intermixing of tuber and functional tissue. This is in agreement with current pathological [4,26] diffusion imaging [27], and with physiological evidence of widespread and variable degrees of pathology in TSC [28–31].

The color coding of the streamline fascicles represents the main direction of the tract: blue for the super-inferior axis, red for the left-right axis, and green for the anterior-posterior axis. The tuber is located at the crosshair, and is partially obscured by the fiber tract, which travels into the tuber tissue. Note that the fibers do not travel through the ventricles; this is an effect from the superimposition of the 3D tracts onto a 2D coronal plane. The full course of the tract is not all in the same plane as the coronal slice.

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