Microelectrode recording findings within the tractography-defined ventral intermediate nucleus

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OBJECTIVE The ventral intermediate nucleus (VIM) of the thalamus is not visible on structural MRI. Therefore, direct VIM targeting methods for stereotactic tremor surgery are desirable. The authors previously described a direct targeting method for visualizing the VIM and its structural connectivity using deterministic tractography. In this combined electrophysiology and imaging study, the authors investigated the electrophysiology within this tractography-defined VIM (T-VIM).

METHODS Thalamic neurons were classified based on their relative location to the T-VIM: dorsal, within, and ventral to the T-VIM. The authors identified the movement-responsive cells (kinesthetic and tremor cells), performed spike analysis (firing rate and burst index), and local field potential analysis (area under the curve for 13–30 Hz). Tremor efficacy in response to microstimulation along the electrode trajectory was also assessed in relation to the T-VIM.

RESULTS Seventy-three cells from a total of 9 microelectrode tracks were included for this analysis. Movement-responsive cells (20 kinesthetic cells and 26 tremor cells) were identified throughout the electrode trajectories. The mean firing rate and burst index of cells (n = 27) within the T-VIM are 18.8 ± 9.8 Hz and 4.5 ± 5.4, respectively. Significant local field potential beta power was identified within the T-VIM (area under the curve for 13–30 Hz = 6.6 ± 7.7) with a trend toward higher beta power in the dorsal T-VIM. The most significant reduction in tremor was also observed in the dorsal T-VIM.

CONCLUSIONS The electrophysiological findings within the VIM thalamus defined by tractography, or T-VIM, correspond with the known microelectrode recording characteristics of the VIM in patients with tremor.

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There is a significant interest in devising direct targeting methods for the thalamic ventral intermediate nucleus (VIM) for tremor surgery.12,25,26 These investigations have in part been driven by 2 major developments: the identification of cerebellar input to the VIM as a therapeutic target26,50 and the introduction of noninvasive procedures (Gamma Knife and MRI-guided focused ultrasound), which dispense with any physiological exploration of the ventral thalamus.15,34,26 We recently described a method for tractography-based VIM (T-VIM) targeting.37 This method involves a step-wise localization of the VIM by checking the anatomical accuracy at each step. First, the borders of the VIM are visualized by tracking the pyramidal tract (PT) and medial lemniscus (ML) (these tracts define the VIM’s lateral and posterior borders, respectively). A VIM region of interest (ROI), similar in dimensions to the human VIM, is then created 3 mm away from both the PT and ML. Finally, the structural connectivity of the VIM with the cerebellum (via the dentatorubrothalamic tract (DRT), which terminates in the VIM) and motor cortex (primary motor and premotor cortex) is confirmed. The reproducibility and clinical application of this methodology was described previously. In this paper we analyzed the electrophysiological findings rela-
tive to T-VIM imaging, within the same patient cohort, to correlate the 2 methods of target localization and further evaluate the merit of imaging findings.

The VIM has a distinct electrophysiological signature in humans. Microelectrode recording (MER) studies have shown that VIM neurons fire within a frequency range of 10 to 15 Hz, exhibit irregular firing patterns, and are often responsive to active movement-responsive (“voluntary” cells), joint movement/pressure-sensitive (“kinesthetic” cells), or active movement- and joint movement/pressure-responsive (“combined” cells). These cells include those that have periodic oscillations in their firing rate at tremor frequency (3–8 Hz), which is frequently coherent with limb or jaw tremor measured using electromyography or an accelerometer. More recently, the local field potential (LFP) within the VIM has been found to have prominent beta oscillations (13–30 Hz), which appears to attenuate when tremor frequency oscillations appear, followed by tremor in the limbs. We hypothesized that these distinct electrophysiological features of the VIM should delineate the borders of this nucleus and therefore coregister with the borders determined by the T-VIM. We studied tremor patients (those with essential tremor and those with tremor-dominant Parkinson’s disease) who underwent preoperative diffusion tensor imaging (DTI) and MER-guided VIM procedures (either thalamotomy or deep brain stimulation [DBS]). The trajectory of the dual recording electrode was analyzed in relation to T-VIM and the findings were categorized into 3 locations—dorsal to, within, and ventral to the T-VIM. We compared the spike pattern (firing rate, burst index), LFP (spectral analysis and area under the curve for the beta oscillations), and tremor efficacy at these 3 different locations.

Methods
The methods used in this study were approved by the University Health Network Research Ethics Board.

Patient Population
Combined tractography and electrophysiology was performed in 6 patients undergoing functional stereotactic procedures for disabling tremor between November 2014 and January 2015. Three patients had essential tremor, and 3 had tremor-dominant Parkinson’s disease. Physiological exploration of the VIM of the thalamus was performed in all patients using the MER-guided technique described elsewhere.

Imaging Protocol
Diffusion tensor imaging (DTI) was performed using a 3-T magnet (GE Signa). The sequence included 60 directions of diffusion gradients (b = 1000 sec/mm$^2$; 0.94 × 0.94 × 3 mm voxel size; TE 86.6 msec; TR 12,000 msec; matrix 128 × 128). 3D fast spoiled gradient echo axial T1-weighted images were also acquired (voxel size 0.85 × 0.85 × 1 mm, matrix 256 × 256, FOV 220 mm, TE 5 msec, TR 12 msec, TI 300 msec). The MR images for intraoperative navigation were obtained with patients placed in a Leksell frame and a 1.5-T magnet (GE Signa Excite). Both volumetric T1- and T2-weighted sequences were obtained at a slice thickness of 1 mm.

VIM Localization on Diffusion Tensor Imaging
The methodology of DTI-based VIM identification has been described in detail previously. Briefly, we used StealthViz software (version 1, Medtronic Inc.) for coregistration of structural and diffusion tensor images and sensor calculation. We then identified the PT and ML, since these tracts form the lateral and posterior boundaries of the VIM nucleus, respectively. A VIM ROI was then created and placed equidistant (3 mm each) from the PT and ML, approximating the axial and sagittal dimensions of the human VIM (4 mm mediolateral, 3 mm anteroposterior, and 6 mm superoinferior). The structural connectivity of this VIM ROI was traced to the contralateral cerebellum and ipsilateral primary motor cortex. For intraoperative guidance, the individual tracts (PT, ML, and VIM fibers) were overlaid on structural T1-weighted images and exported to stereotactic planning software (FrameLink, Medtronic Inc.). The resultant DICOM data set was then rigidly aligned to the preoperative frame-based T1-weighted image used for intraoperative targeting.

Intraoperative Recording and Testing
The VIM was surgically targeted using the standard MER-guided approach. A typical microelectrode trajectory started 10 mm above the intended target and traversed from the anterolateral direction to the posteromedial direction. Each track used 2 microelectrodes (25-µm tip, 600 µm apart, and 0.2-MΩ impedance at 1000 Hz) loaded into a 2-channel cannula and driven forward by 2 independent manual microdrives. The amplified signals were band passed (10–3000 Hz) and simultaneously displayed on an oscilloscope and heard through an audio speaker output. The recordings were also fed to a digital interface (micro1401, Cambridge Electronic Design) for display and saved for offline analysis using Spike2 Neurological Capture (Cambridge Electronic Design). We identified tremor cells and kinesthetic cells and assessed tremor efficacy using clinical stimulations (100 milliamp, 100 Hz) at 1-mm increments. Using this setup, the estimated current spread is typically within 1 mm of the microelectrode tip. The tremor efficacy was categorized into 4 categories based on a comparison of pre- and poststimulation clinical tremor scoring on a 4-point scale: no efficacy (0% reduction or 0 points on the tremor scale), mild efficacy (< 50% reduction or 1 point on the tremor scale), moderate efficacy (≥ 50% reduction or 2 points on the tremor scale), and tremor arrest (100% reduction or > 2 points on the tremor scale). The final electrode implantation was performed at a site with good tremor efficacy and acceptable side effect profile (no lemniscal and pyramidal side effects at clinically relevant stimulation thresholds). All recording data were stored on a central server for offline analysis.

Electrophysiology Analysis
Intraoperative recordings were analyzed offline using Spike2 software (version 7, Cambridge Electronic Design). We identified single-unit activity with a high signal-to-noise ratio (minimum 2:1) and applied a bandpass...
filter (200–3 kHz). Stable recordings (> 10 seconds) that were free from artifacts (noise, speech, motion, and clinical testing as measured by the accelerometer recordings) were isolated. Using the wave-mark function, we isolated spikes using template matching and principal component analysis. Single cells were segregated according to their relation to the VIM, similar to the methodology described previously for the subthalamic nucleus. The length of the VIM trajectory was normalized across subjects on a scale of 0 to 1 (1 corresponding to the top of the T-VIM and 0 corresponding to the bottom of the T-VIM). Thus, the cells ventral to the T-VIM had normalized values less than 0 (negative values), while the cells dorsal to the T-VIM had values greater than 1.

Burst analysis was performed using the Kaneoke and Vitke methodology. Spike events were imported and processed using MATLAB (version 7, Mathworks). The discharge density histograms were then plotted with increasing densities from the left to the right. Burst index was calculated by dividing the mean interspike interval (ISI) by the modal ISI. Other parameters analyzed included the firing rate, mean ISI within the burst, and mean ISI outside of the burst. For the LFP analysis, recording segments (approximately 3 segments per mm) were analyzed along the trajectory. LFP beta power was calculated similar to the methodology described elsewhere. Briefly, recordings were bandpass filtered in the beta frequency range (13–30 Hz) and the beta power of each recording segment was separately calculated by first performing a Fourier transform and then determining the area under the beta frequency peak. This beta power was then plotted against the normalized depth within the VIM as described above.

Statistical Analysis
Data were imported into SPSS (version 22, IBM Corp.) for statistical analysis. Continuous variables were compared using the t-test and ANOVA. Categorical variables were compared using the chi-square test; p < 0.05 was considered statistically significant.

Results
Demographics
The mean age of this cohort was 71.7 ± 5.9 years, and the mean Clinical Rating Scale for Tremor (CRST) subscale B (specific motor tasks/function rating score) was 17.8 ± 5.4. Three patients had essential tremor and 3 had tremor-dominant Parkinson’s disease. In total, 9 microelectrode trajectories were performed. A typical trajectory is shown in Fig. 1.

Distribution of Kinesthetic and Tremor Cells
Overall, 20 kinesthetic and 26 tremor cells were identified (Fig. 2). In relation to the T-VIM, the kinesthetic cells were located mainly in the dorsal part (mean normalized...
depth 0.95 ± 0.49) and the tremor cells in the ventral part (mean normalized depth 0.63 ± 0.44; 2-sample t-test, p = 0.03). The kinesthetic cells responded to movements of the shoulder (2 cells), elbow (14 cells), and wrist (5 cells). Increased firing rate was observed in response to joint movement in all but one instance, in which the thalamic cell was inhibited by elbow flexion.

Spike and Burst Analysis

We identified 27 thalamic cells that met the inclusion criteria for spike analysis. The location of these cells is illustrated in Fig. 2. Overall, 18 cells were localized within the T-VIM, 5 cells were dorsal to the T-VIM, and 4 cells were ventral to the T-VIM. The mean firing rate of cells within the T-VIM was 18.8 ± 9.8 Hz. There was a transi-
tion in the firing rate as the electrode entered the T-VIM. The mean firing rate of thalamic cells located dorsal to the T-VIM was $15.2 \pm 4.4$ Hz compared with $26.6 \pm 10.4$ Hz for the cells within the first few millimeters of electrode entry into the T-VIM (2 sample t-test, $p = 0.2$). As shown in Fig. 2 we noticed a higher amplitude of action potential within the T-VIM, suggesting large neurons. Both the cells within and outside the T-VIM exhibited the typical irregular firing pattern with burst indices of $4.5 \pm 5.4$ and $3.8 \pm 0.8$, respectively. However, there was a significant association between burst index and depth within T-VIM (1-way ANOVA, $p < 0.001$), with cells recorded toward the bottom of T-VIM tending to have a higher burst index.

**LFP Analysis**

Significant LFP beta power was identified within the T-VIM (area under the curve for 13–30 Hz = $6.6 \pm 7.7$). There was a trend toward higher beta power in the dorsal T-VIM (Pearson’s correlation coefficient $R^2$) between beta power and normalized depth = $0.35$, $p = 0.08$). A representative track from one patient is reconstructed for the area under the main spectral peak for the 13- to 30-Hz frequency (Fig. 2).

**Tremor Efficacy**

Microelectrode stimulation within the T-VIM resulted in significant reduction in tremor ($50\% \pm 34.9\%$ corresponding to a 2-point difference in the tremor rating score). Although stimulation reduced tremor outside the T-VIM, the magnitude of this reduction was smaller ($30\% \pm 29.4\%$ decrease in tremor score; 2-sample t-test, $p = 0.08$). The most significant reduction in tremor was observed in the dorsal T-VIM ($64\% \pm 19.5\%$ reduction in dorsal T-VIM vs $25\% \pm 17.3\%$ reduction immediately above the VIM, 2-sample t-test, $p = 0.02$).

**Clinical Outcomes**

The $x$, $y$, and $z$ coordinates of the T-VIM and surgically identified VIM in relation to the posterior commissure were $14.3$, $7.2$, and $0.3$ and $13$, $6.5$, and $0.5$, respectively.

Over the long term (follow-up 8.9 ± 2.6 months, range 5–11 months), the CRST subscale B improved by 56% (SD $11.9\%$, range 41%–75%). The CRST subscale B improvement on the operated side improved by 78% (SD $17.2\%$, range 50%–100%).

**Discussion**

In this combined imaging and electrophysiology study, we established that the electrophysiological findings within the area of the thalamus defined by tractography or T-VIM confirm the known MER characteristics of the VIM in patients with tremor.

At present, refractory essential tremor is most often surgically treated by targeting the VIM with radiofrequency lesioning or DBS.\(^{39}\)\(^{39}\)\(^{39}\) Gamma Knife lesioning,\(^{36}\) and more recently with MR-guided focused ultrasound lesioning.\(^{34}\)\(^{34}\) While the outcome of these treatments is crucially dependent on precise VIM localization,\(^{36}\) stereotactic VIM targeting relies on indirect methods because the structural MR images (both 1.5 and 3 T) fail to visualize individual thalamic nuclei.\(^{9}\)\(^{18}\) Indirect targeting relies on standard coordinates without accounting for the interindividual variability in thalamic structure and location.\(^{10}\)\(^{23}\)\(^{25}\)

This may potentially result in heterogeneous outcomes after tremor surgery.\(^{5}\)\(^{16}\) More recently, emerging noninvasive techniques\(^{43}\)\(^{54}\)\(^{56}\) preclude the confirmation with MER mapping, creating an immediate need for reliable, direct targeting of the VIM. Several techniques have previously been investigated, including gradient-filtered T1-weighted imaging,\(^{11}\) T1 and T2 dependence-weighted 3-T imaging,\(^{46}\) multiparametric (“virtual-dot-com imaging”) 3-T imaging,\(^{55}\) optimized magnetization transfer mapping with 3-T imaging,\(^{17}\) susceptibility-enhanced T1-weighted 3T imaging,\(^{34}\) and susceptibility-weighted 7-T imaging.\(^{1}\) However, so far, none of these have been translated into clinical application.

A dysfunction within the cerebellothalamocortical network has recently been implicated in tremor pathophysiology.\(^{7}\)\(^{13}\)\(^{14}\)\(^{35}\)\(^{42}\)\(^{44}\)\(^{48}\) Indeed, functional neuroimaging studies associate the successful modulation of this network with good surgical outcomes after VIM DBS\(^{26}\) and thalamotomy.\(^{50}\) In particular, the effective DBS contacts appear to lie close to the DRT.\(^{12}\)\(^{25}\) The emerging direct targeting methods for the VIM have also adopted this network-centered approach. Some studies have demarcated thalamic subregions based on their inherent diffusion characteristics\(^{47}\)\(^{49}\)\(^{57}\) or their connectivity with different cortical regions,\(^{8}\)\(^{45}\) and more recently have used probabilistic\(^{22}\)\(^{23}\)\(^{28}\)\(^{40}\) and deterministic\(^{3}\)\(^{12}\)\(^{25}\)\(^{52}\)\(^{53}\) DTI algorithms to directly map the DRT. While novel and valuable, these approaches have not found a clinical application due to several limitations; for example, they are often computationally intensive, poorly integrated with stereotactic targeting platforms, or too imprecise for clinical application, where an accuracy of less than 1 mm is desired.\(^{43}\) We recently described a tractography-based targeting method that identified the T-VIM according to the relative location of the surrounding PT and ML, which are easily visualized with DTI.\(^{37}\) Selecting a thalamic subregion of comparable size to the anatomical VIM,\(^{20}\) we confirmed the structural connectivity of the T-VIM with the cerebellum and motor cortex. Overall, we found this method to be accurate, reproducible, and useful for stereotactic VIM targeting. Notably, we found a good correlation between the coordinates of T-VIM and the final surgical VIM target, implying that the T-VIM is in the vicinity of the actual VIM.

In this investigation, we analyzed MER data to interrogate the electrophysiological properties of the T-VIM and thereby understand its relationship to the anatomical VIM. We found that the cells recorded within the T-VIM have firing rate and burst characteristics similar to those seen in the human VIM.\(^{29}\)\(^{31}\) In further concordance with the literature, we also identified kinesthetic cells\(^{30}\)\(^{32}\) and tremor cells (with a typical oscillatory frequency of around 6 Hz, and corresponding to tremor)\(^{31}\)\(^{33}\) throughout the electrode tract often located above, within, and below the T-VIM. The LFP analysis also yielded results in line with other published reports,\(^{6}\) as we found significant beta oscillatory activity within the T-VIM. Finally, microelectrode stimulations confirmed that the zone of greatest tremor efficacy was co-localized.

1. Microelectrode recording VIM tractography
2. J Neurosurg Volume 126 • May 2017
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with the T-VIM. There is a lack of clarity in the published imaging literature about whether DRT terminations truly represent the anatomical VIM or whether it instead represents other neighboring structures like thalamic nucleus ventralis oralis posterior, caudal zona incerta, and posterior subthalamic region.\textsuperscript{12,38} Our findings in this combined imaging and electrophysiology study suggest that DRT terminations within the T-VIM do in fact represent the VIM subregion of the thalamus.

Several limitations of this study should be noted. First, we have used a single tensor tractography algorithm for delineating the T-VIM that has poor resolution for crossing fibers. However, since the VIM is a relay nucleus,\textsuperscript{19} the better resolution afforded by more advanced algorithms\textsuperscript{8,43} does not offer a significant advantage in visualizing the T-VIM. The algorithm used for this study was chosen due to its ease of tensor calculation, tract visualization, and integration with the stereotactic targeting platform. Another limitation was the small sample size of this study, which precluded us from describing either the somatotopy or tremor efficacy zones within the T-VIM.

Conclusions

The electrophysiological findings within the VIM defined by tractography, or T-VIM, correspond with the known MER characteristics of the VIM. In the future, long-term tremor outcomes associated with T-VIM targeting and their comparisons with conventional MER-guided methods are desirable.

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Disclosures

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

Author Contributions

Conception and design: Krishna, King. Acquisition of data: Krishna, King, Basha, Elias, Sammartino. Analysis and interpretation of data: Krishna, King, Basha, Elias, Hutchison. Drafting the article: Krishna, Basha. Critically revising the article: Sameen M, et al. All authors approved the final version of this manuscript. All authors agreed to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. An equally contributing author is listed as such.

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