Dorsolateral subthalamic neuronal activity enhanced by median nerve stimulation characterizes Parkinson’s disease during deep brain stimulation with general anesthesia

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OBJECT Deep brain stimulation (DBS) surgery under general anesthesia is an alternative option for patients with Parkinson's disease (PD). However, few studies are available that report whether neuronal firing can be accurately recorded during this condition. In this study the authors attempted to characterize the neuronal activity of the subthalamic nucleus (STN) and elucidate the influence of general anesthetics on neurons during DBS surgery in patients with PD. The benefit of median nerve stimulation (MNS) for localization of the dorsolateral subterritory of the STN, which is involved in sensorimotor function, was explored.

METHODS Eight patients with PD were anesthetized with desflurane and underwent contralateral MNS at the wrist during microelectrode recording of the STN. The authors analyzed the spiking patterns and power spectral density (PSD) of the background activity along each penetration track and determined the spatial correlation to the target location, estimated using standard neurophysiological procedures.

RESULTS The dorsolateral STN spiking pattern showed a more prominent bursting pattern without MNS and more oscillation with MNS. In terms of the neural oscillation of the background activity, beta-band oscillation dominated within the sensorimotor STN and showed significantly more PSD during MNS (p < 0.05).

CONCLUSIONS Neuronal firing within the STN could be accurately identified and differentiated when patients with PD received general anesthetics. Median nerve stimulation can enhance the neural activity in beta-band oscillations, which can be used as an index to ensure optimal electrode placement via successfully tracked dorsolateral STN topography.

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KEY WORDS deep brain stimulation; general anesthesia; median nerve stimulation; microelectrode recording; subthalamic nucleus; functional neurosurgery
Microelectrode recording (MER) is possible when patients are deeply sedated, such as during general anesthesia. Microelectrode mapping during STN-DBS is an essential procedure to optimize the placement of DBS electrodes along the STN. The somatotopic organization of the STN was determined using MER, which showed that the movement-related cells that control the arm were located laterally and at the rostral and caudal poles of the STN, whereas the cells that control leg movement were located medially and centrally. It is crucial to implant the DBS electrode into this motor subregion to ensure a good outcome. Although somatotopic mapping within the STN during MER may facilitate optimal placement of the DBS electrodes, the associated shortcomings of this process are that it is time consuming and requires the patient to be awake during the procedure in most hospitals. In our previous study, we showed the feasibility of conducting MER during conditioned general anesthesia under desflurane, which allowed a good long-term outcome. However, the topography of the STN may not be delineated by MER signals due to partial anesthetic suppression. This characteristic may lead to a higher stimulation side effect during clinical follow-up. In addition to determining the correlation between the STN topography and the characteristic signal, enhancing MER neuronal signals under conditioned general anesthesia might facilitate the improvement of DBS outcomes.

The pathological and characteristic neuronal oscillations of the STN have been revealed in animal models of PD and human patients. Beta-band (13–30 Hz) oscillations are not only distinct to the dorsolateral STN and able to be used to predict DBS electrode placement, but can also be manipulated by a patient’s active movement. Placing the stimulating electrode on “target” is the crucial factor for a favorable outcome. In addition to movement-related change of STN firing, previous studies have also tried to localize the target area of the basal ganglia nuclei and STN using somatosensory evoked potentials (SSEPs) elicited by median nerve stimulation (MNS). However, SSEPs coupled with MNS failed to demonstrate the characteristic changes on STN regions, and these studies used macroelectrode recordings. Furthermore, neurophysiological recordings have been widely employed recently to investigate the status of cortical neurons and altered levels of unconsciousness under general anesthesia. By examining conditioned general anesthesia during surgery, this protocol may allow us to study the extent to which general anesthesia can modulate neuronal firing in the STN. The application of MNS may also explain the mechanisms by which peripheral nerve stimulation may facilitate or hinder CNS oscillation.

Methods

Patient Selection

Eight consecutive patients with PD who underwent bilateral STN-DBS in the Hualien Tzu Chi General Hospital from January 2010 to November 2012 were enrolled in this study (Table 1). All patients met the United Kingdom PD Brain Bank diagnostic criteria, with at least 2 of the cardinal symptoms present. Before surgery, each patient underwent a levodopa test to confirm a positive levodopa response (> 28% improvement in the Unified Parkinson’s Disease Rating Scale [UPDRS] Part III score). Brain MRI was performed preoperatively to rule out structural abnormalities in each patient. All medications were withdrawn at least 12 hours before surgery. Before surgery in the medication “off” condition, the mean Hoehn and Yahr score was 3.0 ± 0.46 (range 2.5–4) and the mean UPDRS-III score was 41.1 ± 10.3 (range 27–57). The Institutional Review Board at Tzu Chi General Hospital approved the surgical and evaluation procedures. Informed consent was obtained from each patient.

Imaging and Targeting

Images were obtained using a 1.5-T MRI unit (General Electric). The standard settings included T1-weighted axial images of 0.75-mm thickness and T2-weighted axial images of 2-mm thickness. Each sequence was performed in contiguous slices. The images were transferred to the DICOM database using the Picture Archiving and Communication System and the stealth neuronavigation workstation (Medtronic). The image fusion software fused the two sets of MR images to form a 3D reconstruction. The tentative surgical target coordinates for the tip of the permanent implantable electrode were set at the central, lowest border of the STN by direct visualization on MRI, as previously described in detail. A Leksell G-frame unit (Elekta Instrument, Inc.) was used for the stereotactic procedure. The patient was resting in a straight supine position, and the head frame was secured in a Mayfield adapter. The target coordinates were applied to the stereotactic frame and the working stage.

Anesthetic Procedure

All patients received a general anesthetic with endotracheal intubation. Anesthesia was induced initially by administering regular narcotic agents. All patients were maintained using desflurane inhalation during the surgical procedures. The depth of the anesthesia was maintained at 0.5–1.0 minimal alveolar concentration (MAC), and each patient’s heart rate and blood pressure were monitored to ensure that the patient avoided a cough reflex or any change in heart rate and blood pressure during the MER procedure.

Electrical Stimulation of the Median Nerve

A Digitimer constant current stimulator (model DS7A) was used to apply electrical stimulation to the contralateral median nerve. A stimulation electrode was placed on the wrist (cathode, median nerve 2 cm proximal to the wrist crease; anode, 2 cm distal to the cathode), and the stimulation parameters included a pulse width of 0.2 μsec, an intensity of 30 mA, and a frequency of 33 Hz.

Microelectrode Recording Procedure

The MER device is 10–40 μm in diameter and measures 200 mm in length, with a 10-mm-long bare tungsten tip (FHC). The recording impedance was usually between 0.5 and 1 MΩ. The microelectrode was mounted on a
were permanently implanted. Electrode insertion tests after the electrodes (Medtronic 3387 DBS leads) were used to localize the STN neuron. We did not perform any macrostimulation tests after the electrodes were inserted. Passive movements related to the activity of the STN were tested during MER to determine whether any movement-related neuronal firing changes occurred. Neurons were regarded as movement related if there was an audible alteration in the neuronal discharge that was reproducible and synchronous with the passive movement of the contralateral limbs. The analysis criteria for MERs included spike detection, spike pattern differentiation, neural background activity extraction, and power spectral density (PSD) estimation.

**Spike Detection**

Neuronal activity recordings of the dorsolateral STN were exported offline to the Spike 2 software suite (Version 5, Cambridge Electronic Device). In addition, the sampling rate of all MERs was 24 kHz. Spikes were differentiated for spike detection using threshold detection and template matching. Identified single units were verified using principle component analysis and visual inspection (Fig. 1). However, setting the proper threshold was critical for spike detection.

Typically, the MER signal consists of the spiking activity of most nearby neuronal units and background electrical activity due to several sources, including more remotely located units, measurement noise, and various artifacts. In this study, we assumed the input signal \( x(t) \) to be a linear combination of transient action potential signals (\( x_{ap} \)) and band-limited Gaussian noise (\( x_n \)), which is shown as follows:

\[
x(t) = x_{ap} + x_n
\]

Moreover, we assumed that the Gaussian background noise had a mean of zero. In other words, the noise can be fully described by its root mean square value, which is equivalent to its standard deviation \( \sigma \). In this study, the detection threshold was set at 3 times the estimated background root mean square. When the waveform amplitude exceeded the threshold, we searched for the signal

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Age of Onset (yrs), Sex</th>
<th>Disease Duration (yrs)</th>
<th>Hoehn &amp; Yahr Stage (off)</th>
<th>Levodopa Response in UPDRS Part III</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Drug Off</td>
</tr>
<tr>
<td>1</td>
<td>53, M</td>
<td>7</td>
<td>2.5</td>
<td>38</td>
</tr>
<tr>
<td>2</td>
<td>44, F</td>
<td>12</td>
<td>3</td>
<td>45</td>
</tr>
<tr>
<td>3</td>
<td>59, M</td>
<td>13</td>
<td>3</td>
<td>32</td>
</tr>
<tr>
<td>4</td>
<td>52, F</td>
<td>8</td>
<td>3</td>
<td>57</td>
</tr>
<tr>
<td>5</td>
<td>52, F</td>
<td>10</td>
<td>3</td>
<td>39</td>
</tr>
<tr>
<td>6</td>
<td>37, M</td>
<td>8</td>
<td>3</td>
<td>54</td>
</tr>
<tr>
<td>7</td>
<td>53, M</td>
<td>11</td>
<td>3</td>
<td>27</td>
</tr>
<tr>
<td>8</td>
<td>48, F</td>
<td>8</td>
<td>2.5</td>
<td>37</td>
</tr>
</tbody>
</table>

Mean ± SD: 49.8 ± 6.7, 9.6 ± 2.2, 3.0 ± 0.5, 41.1 ± 10.3, 20.0 ± 6.1, 50.0 ± 15.7

*Calculated by: drug off – drug on/drug off.
wave for a distinct spike. The threshold setting was calculated using Matlab (MathWorks).

Spike Pattern Differentiation

The spike times of identified single units were imported into Neuroexplorer (Version 4, Nex Technologies) to determine the mean firing rates, interspike intervals (ISIs), and autocorrelograms. The asymmetry index and the coefficient of variance of the ISI were calculated. Autocorrelograms of the single-unit firing patterns were characterized into 3 categories: 1) irregular, which exhibited an initial trough that rose smoothly to a steady state; 2) bursty, which was characterized by an initial peak followed by a decay to a steady state; and 3) oscillatory, in which discharges with similar refractory periods exhibited repetitive neuronal firing with multiple equidistant peaks and troughs (Fig. 2).22

Neural Background Activity Extraction and PSD Estimation

Background activity extraction is an essential step while examining the neuronal background activities of the dorsolateral STN evoked by MNS. In this process, the traces in the segments from 0.5 μsec before to 2.5 μsec after each spike timestamp were replaced by a random spike-free 3-μsec consecutive signal from a random location within the same trace.17 Moreover, to explore the oscillatory characteristics of the background activity, we determined the PSD of beta-band activity (13–30 Hz). The PSD was estimated using Welch’s method. The signal was divided equally into segments with 50% overlap. Each section was windowed with Hamming windows whose length was equal to the number of samples in 2 seconds.

Statistical Analysis

Numerical values of clinical outcomes measured before and after the operation were compared using paired t-tests. The firing rate, asymmetry index, and coefficient of variance of each ISI were compared between the MSN on and off conditions using paired t-tests. The proportions of various firing patterns with MNS either on or off were compared using chi-square tests; p values < 0.05 were considered statistically significant.

Results

Clinical Outcome After STN-DBS and Localization

Postoperative DBS substantially improved the clinical status of the patients. A follow-up examination at 3 months after surgery revealed that STN-DBS significantly improved UPDRS-III scores by 38.68% (med off and DBS off, 38.0 ± 3.61; med on and DBS on, 23.3 ± 5.86; p < 0.001). The mean stimulation parameters of channel 1 (left STN-DBS electrode)/channel 2 (right STN-DBS electrode) were 3.23 ± 0.23 V and 2.91 ± 0.70 V, respectively; 60-μsec (both channels) pulse width; and frequencies of 122.5 ± 13.89 Hz and 118.75 ± 15.53 Hz, respectively.

Analysis of Single Dorsolateral STN Unit Activity

We performed a mean of 57.13 ± 9.19 MNS stimulations along the STN path. The mean number of MER trajectories was 1.33 ± 0.82 per patient side and the recorded length of the STN was 4.69 ± 0.82 mm. The mean firing frequencies for 10-second intervals of the MNS off and on conditions were 34.40 ± 26.16 Hz (n = 46) and 30.85 ± 20.25 Hz (n = 66), respectively, which were not significantly different (p > 0.05, t-test). The asymmetry index and coefficient of variance of the ISIs before and after MNS were similar (Table 2). Based on the autocorrelogram analysis, the most frequently observed activity pattern of all dorsolateral STN units with MNS off was bursty (19/46 units, 41.3%), followed by irregular (16/46 units, 34.7%) and oscillatory (11/46 units, 24.0%) firing patterns. Moreover, the observed activity patterns of all dorsolateral STN units with MNS on were oscillatory (31/66 units, 47.0%), followed by bursty (19/66 units, 28.8%) and irregular (16/66 units, 24.2%) firing patterns. Oscillatory neuronal activity was found with a significantly increased incidence between MNS off and on (odds ratio 2.82, p < 0.05, chi-square test). In contrast, the bursty firing patterns were significantly decreased in MNS on relative to MNS off (odds ratio 0.57, p < 0.05, chi-square test).

Analysis of Background Oscillation

Oscillatory activity of the neuronal firing pattern is correlated with the synchronous background activity of the MER data. To explore the background oscillation, we estimated the spectrum character (i.e., the PSD) of the background activity while neuronal spikes were detected. In addition, the PSD of dorsolateral subthalamic neurons in the beta band exhibits significant features. In this study, the PSD of background activity in the beta band was examined to explore the oscillation evoked by MNS. Both the PSD of each patient and that of the whole group in the
beta band were significantly increased while MNS was on compared with while MNS was off (Table 3; t-test).

**Discussion**

After we analyzed the physiological features of the STN from single neuron recordings in patients with PD, we demonstrated that STN neurons could be recorded and identified while patients were under general anesthesia. In addition to discriminating the characteristics among 3 types of STN firing while a patient was under general anesthesia, we could identify pattern shifts of STN neuronal firing after peripheral nerve stimulation using MNS.

A clear distinction of the anatomical boundaries among the 3 subregions (motor, associative, and limbic areas) of the STN is difficult to achieve in human subjects. Through strict mapping of the MER axis and preoperative MRI, we determined the characteristics of the dorsolateral and motor subterritory neurons of the STN in patients with PD under general anesthesia, which were mostly in

**TABLE 2.** The firing frequency, as well as the coefficient of variance and asymmetry index of the ISI, within the dorsolateral STN

<table>
<thead>
<tr>
<th>MNS Spike Pattern</th>
<th>No. of Neurons</th>
<th>%</th>
<th>Firing Frequency (Hz)</th>
<th>Coefficient of Variance</th>
<th>ISI Coefficient of Variance</th>
<th>Asymmetry index</th>
</tr>
</thead>
<tbody>
<tr>
<td>Off</td>
<td>Irregular</td>
<td>16</td>
<td>34.7</td>
<td>58.28 ± 29.41</td>
<td>1.04 ± 0.19</td>
<td>0.34 ± 0.15</td>
</tr>
<tr>
<td></td>
<td>Bursty</td>
<td>19</td>
<td>41.3</td>
<td>25.43 ± 12.21</td>
<td>1.36 ± 0.40</td>
<td>0.25 ± 0.12</td>
</tr>
<tr>
<td></td>
<td>Oscillatory</td>
<td>11</td>
<td>24.0</td>
<td>16.37 ± 8.42</td>
<td>1.33 ± 0.21</td>
<td>0.13 ± 0.04</td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>46</td>
<td>100</td>
<td>34.40 ± 26.16</td>
<td>1.22 ± 0.33</td>
<td>0.25 ± 0.15</td>
</tr>
<tr>
<td>On</td>
<td>Irregular</td>
<td>16</td>
<td>24.2</td>
<td>42.69 ± 26.97</td>
<td>0.87 ± 0.19</td>
<td>0.50 ± 0.32</td>
</tr>
<tr>
<td></td>
<td>Bursty</td>
<td>19</td>
<td>28.8</td>
<td>36.77 ± 20.43</td>
<td>1.26 ± 0.33</td>
<td>0.24 ± 0.08</td>
</tr>
<tr>
<td></td>
<td>Oscillatory</td>
<td>31</td>
<td>47.0</td>
<td>21.11 ± 9.06</td>
<td>1.27 ± 0.35</td>
<td>0.16 ± 0.06</td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>66</td>
<td>100</td>
<td>30.85 ± 20.25</td>
<td>1.17 ± 0.36</td>
<td>0.27 ± 0.22</td>
</tr>
</tbody>
</table>
agreement with previously published results in terms of the asymmetry index and coefficient of variance of the ISIs. It is interesting that lower mean firing frequencies (34.4 Hz) were recorded in patients under general anesthesia compared with patients undergoing DBS with a local anesthetic (41.2–58 Hz).22,23 It has been debated whether it is possible to record STN firing while patients receive general anesthesia and lose consciousness, as well as how these agents, such as intravenous propofol or inhaled desflurane, might dampen neuronal activity. The findings of our study suggest that performing meticulous monitoring while administering desflurane to implant DBS electrodes in patients with PD not only allows successful electrophysiological mapping but also avoids precluding a subset of patients who are not suitable for surgery under local anesthesia.8

The dorsolateral subterritory of the STN has long been considered an important part of the sensorimotor circuit of the basal ganglio-thalamo-cortical network and implicated as underpinning the pathophysiology of PD.1,10,11 In accordance with clinical observations, high-frequency DBS within this region has also been demonstrated to achieve cordance with clinical observations, high-frequency DBS oscillations.7 Sensory stimulation could change motor cortex excitability, and a sensory-conditioned stimulus such as MNS could inhibit the motor cortex, which may indirectly lead to the increased oscillation of STN neurons.21

Additionally, MNS has been used as a reliable tool to identify basal ganglia structures from SSEP recordings and has been even more widely applied toward neural firing characterization and elucidation of the mechanisms of neural oscillation. Most of these reports chose local field potential (LFP) to identify the specific features of STN,12,18 In the other reports, using single neuronal discharges to facilitate delineation of STN in patients with PD has also been proven to be feasible intraoperatively or through offline analysis.17,21 Another study that analyzed the relationship between LFP and single neuronal discharges in STN showed that LFP illustrated synchronous activity of the neuronal population.13 Taking all of these data together, it is very difficult to decide which neuronal cacophony is better, but both LFP and single neuronal discharge play pivotal roles for us to treat disease with neurophysiological problems, such as PD.

Compared with STN firing without MNS, the mean firing frequencies of STN neurons, as well as the asymmetry index and coefficient of variance of the ISIs, remained similar during MNS. Nonetheless, our observation that the PSD of beta-band firing—which has been evaluated further during dorsolateral STN analysis—was remarkably augmented during MNS suggests that MNS can also enhance low-frequency oscillation in addition to ultra-fast-frequency oscillation, and facilitate single-unit recording of STN neurons during general anesthesia.7

Although we can achieve recording of the STN under general anesthesia, we did encounter several difficulties that may explain why the mean firing rate was lower under desflurane general anesthesia. First, prolonged surgical duration may cause an accumulation of anesthetic concentration. In such circumstances, we need to lower the MAC of the patient and wait for a proper regain of signal. Second, a damaged MER electrode may hinder accurate neural signal recording. We need to check the impedance of the microelectrode regularly during MER. Lastly, we relied on the MAC to ensure proper sedation. However, we found a nonlinear correlation between the level of the MAC and characteristics of MER. Whether other physiological signatures could help us quantify the depth of anesthesia more accurately may need further research.

**Conclusions**

Analyzing the characteristics of MER within the dorsolateral STN during baseline and MNS confirmed the feasibility of differentiating STN firing while patients with PD undergo surgery under general anesthesia. MNS not only allows the enhancement of background oscillations and changes in spiking patterns but also may facilitate localization of the STN for patients with PD under general anesthesia and allow more candidates to undergo DBS operations.

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References


Author Contributions

Acquisition of data: Tsai, Chuang. Analysis and interpretation of data: Tsai, Chuang, Kuo. Drafting the article: Tsai, Chuang. Critically revising the article: SY Chen. Reviewed submitted version of manuscript: SY Chen, Tsai, Kuo, Chao, Hung. Approved the final version of the manuscript on behalf of all authors: SY Chen. Administrative/technical/material support: YT Chen.

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