The Medial Subthalamic Nucleus Border as a New Anatomical Reference in Stereotactic Neurosurgery for Parkinson’s Disease

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Abstract

Introduction: The intersection of Bejjani’s line with the well-delineated medial subthalamic nucleus (STN) border on MRI has recently been proposed as an individualized reference in subthalamic deep brain stimulation (DBS) surgery for Parkinson’s disease (PD). We, therefore, aimed to investigate the applicability across centers of the medial STN border as a patient-specific reference point in STN DBS for PD and explore anatomical variability between left and right mesencephalic area within patients. Furthermore, we aim to evaluate a recently defined theoretic stimulation “hotspot” in a different center. Methods: Preoperative 3-Tesla T2 and susceptibility-weighted images (SWI) were used to identify the intersection of Bejjani’s line with the medial STN border in left and right mesencephalic area. The average stereotactic coordinates of the center of stimulation relative to the medial STN border on T2 sequences were 3.1 mm lateral, 0.7 mm anterior, and 1.8 mm superior, in proximity of the predefined theoretic stimulation “hotspot.” Results: Fifty-four patients provided 108 stereotactic coordinates of medial STN borders on both sequences. Significant difference in means was found in the Y-(anteroposterior) and Z-(dorsoventral) directions (T2 vs. SWI; \( p < 0.001 \)). Mean coordinates in the Y-(anteroposterior) direction differed significantly between left and right mesencephalic area (T2: \( p < 0.001 \); SWI: \( p = 0.021 \)). Sixty-six DBS leads were placed in 36 patients that had finished stimulation programming, and the average stereotactic coordinates of the center of stimulation relative to the medial STN border on T2 sequences were 3.1 mm lateral, 0.7 mm anterior, and 1.8 mm superior, in proximity of the predefined theoretic stimulation “hotspot.” Conclusion: The medial STN border is applicable across centers as a reference point for STN DBS surgery for PD and seems suitable in order to account for interindividual and intraindividual anatomical variability if one is aware of the discrepancies between T2-weighted imaging and SWI.

Keywords
Deep brain stimulation · Parkinson’s disease · Subthalamic nucleus · Red nucleus · T2-weighted imaging · Susceptibility-weighted imaging

Introduction

Deep brain stimulation (DBS) of the dorsolateral sensorimotor part of the subthalamic nucleus (STN) is considered the most effective surgical treatment for medically refractory Parkinson’s disease (PD) [1]. However,
clinical outcome after STN DBS varies widely between patients [2–5]. In part, this may be explained by individual anatomical variability of the STN combined with poor visualization of the dorsolateral sensorimotor part of the STN with current imaging techniques [6, 7].

The midcommissural point (MCP) is currently the anatomical reference point in stereotactic neurosurgery for STN DBS. By identifying the anterior commissure (AC) and posterior commissure (PC) on MRI, stereotactic coordinates relative to the MCP are used for preoperative planning, intraoperative targeting, and DBS lead placement, and postoperative localization of the active DBS lead contacts. The interpretation of stereotactic coordinates without accounting for patient’s anatomical variability, however, is of questionable merit [8–12].

In a recent publication, Bot et al. [12] proposed the medial STN border (defined as the intersection of Bejjani’s line [13] with the medial border of the STN) as a new, individualized, reference point that is well delineated on standard MR imaging [12]. In their study, the authors found a significant correlation between clinical improvement and the distance to the active stimulation contacts from this medial STN reference point; in contrast, such a correlation was not found between clinical improvement and distance to active stimulation contacts from the MCP. The authors went on to define a theoretic stimulation “hotspot” within the STN, at 2.8 mm lateral, 1.7 mm anterior, and 2.5 mm superior to the medial STN border. More recently, using the same methods in a second, larger patient cohort, this theoretic stimulation “hotspot” was refined and pinpointed to 2.6 mm lateral, 0.7 mm anterior, and 1.9 mm superior to the medial STN border on T2-weighted imaging (throughout this manuscript referred to as “predefined theoretic stimulation ‘hotspot’”; Bot et al. Deep brain stimulation for Parkinson’s disease: refining the optimal location within the subthalamic nucleus. Unpublished data and abstract presentation at XXIIIrd Congress of the European Society of Stereotactic and Functional Neurosurgery, Edinburgh, UK, September 28, 2018). Again, the medial STN border proved superior compared to the MCP as anatomical reference for correlation of DBS location and motor improvement. However, the applicability of the medial STN border as a reference point in DBS surgery for PD across different centers remains unknown.

As acknowledged by the authors, the mean stereotactic coordinates of the medial STN border (N = 65, 3-Tesla T2-weighted images) varied considerably, at 9.2 ± 1.1 (range 6.8–11.5) mm lateral, 2.8 ± 0.8 (range 1.5–5.3) mm posterior, and 4.2 ± 1.0 (range 2.0–6.1) mm inferior to the MCP [12]. This supports the need to account for interindividual anatomical variability in STN DBS surgery. In addition, morphometric variability of the STN and the red nucleus (RN) between left and right mesencephalic area within patients with advanced PD has been reported [14]. This suggested intraindividual anatomical variability could be relevant in localizing the medial STN border in DBS surgery for PD.

As susceptibility-weighted imaging (SWI) gained interest in DBS surgery for PD due to its better visualization of the STN compared to T2-weighted imaging, discrepancies between T2-weighted imaging and SWI have been a subject of study [6, 7, 15, 16]. In localizing the medial STN border, the correspondence between T2-weighted imaging and SWI remains unclear. When employing the medial STN border as an individualized anatomical reference point in DBS surgery for PD, however, this information is of value.

In the current study, we aimed to investigate the applicability across centers of the medial STN border as a new, individual reference point on both 3-Tesla T2-weighted imaging and SWI, and explored anatomical variability between left and right mesencephalic area within patients. In addition, we aimed to evaluate the recently refined theoretic stimulation “hotspot” in PD patients that underwent STN DBS surgery in a different center.

**Materials and Methods**

**Patient Selection and Clinical Data Collection**

We selected patients with PD who underwent STN DBS at Rush University Medical Center (RUMC) between October 2016 and August 2018. We reviewed medical charts and preoperative imaging used for the stereotactic procedure. Patients were included in the current analysis if preoperative MR imaging included both T2-weighted and susceptibility-weighted (SWI) sequences. Patients with a history of any intracranial surgery were excluded. We extracted clinical data comprising age, gender, disease duration (defined as years between symptom onset and date of surgery), preoperative levodopa responsiveness (change in UPDRS-III scores between OFF and ON medication), the levodopa-equivalent daily dose (LEDD) [17] at baseline and follow-up, and active DBS lead contact(s).

**Surgical Procedure**

All patients underwent awake, single-track MER-guided, frame-based DBS lead placement under local anesthesia. One to 2 weeks prior to the day of surgery, patients underwent MRI (Magnetom Verio 3-Tesla, Siemens, Munich, Germany). Post-contrast 3D volumetric T1-weighted imaging, T2-weighted imaging, and SWI were merged, and target planning was performed using the StealthStation S7 equipped with Framelink software (Medtronic,
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The Medial STN Border

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Stimulation, the midpoint between active contacts was set as center

neuronally, for instance in double monopolar, bipolar, or interleaving

quisition time 11:40 min.

20 ms; slice thickness 1 mm; voxel size 0.5 × 0.5 × 1.0 mm; and ac-

the parameters were as follows: repetition time 30 ms; echo time

el size 0.7 × 0.7 × 2.0 mm; and acquisition time 8:47 min. For SWI,

level of 280 and a width of 264 were chosen for all images.

of 1,080–1,180 was chosen for T2-weighted images. For SWI, a

ation window to depict STN borders a level of 820 and a width range

on 3-Tesla T2-weighted sequences and SWI. For optimal resolu-

 reviewers (EB and LVM) for both left and right mesencephalic area

depicted the RN at its maximum diameter, the plane closest to 4

der of the RN, perpendicular to the AC-PC line (Bejjani’s line

crosshair was used to draw a line coinciding with the anterior bor-

plane in which the RN was visualized at its maximum diameter,

we identified the medial STN border by determining the axial

1 week after DBS lead placement.

lead placement. Generally, implantable pulse generators were

thresholds for therapeutic effect and side effects. In each hemi-

esthetic cells [8, 21–24] and performing test stimulation to assess

refine targeting by identifying the electrophysiological dorsal and

ventral STN border [20], assessing for presence or absence of kin-

esthetic cells [8, 21–24] and performing test stimulation to assess

accuracy of the first microelectrode track, as well as final DBS

lead placement. Generally, implantable pulse generators were

placed 1 week after DBS lead placement.

Identification of the Medial STN Border

According to methods previously described by Bot et al. [12],

we identified the medial STN border by determining the axial

plane in which the RN was visualized at its maximum diameter,

using both axial- and coronal-orientated imaging. In this plane, a

crosshair was used to draw a line coinciding with the anterior bor-

der of the RN, perpendicular to the AC-PC line (Bejjani’s line

[13]). The point of intersection with the STN was localized and

stereotactic coordinates relative to the MCP were retrieved and

defined as the medial STN border (Fig. 1). If more than 1 plane

depicted the RN at its maximum diameter, the plane closest to 4

mm below MCP was chosen. This was done by consensus from 2

reviewers (EB and LVM) for both left and right mesencephalic area

on 3-Tesla T2-weighted sequences and SWI. For optimal resolu-

tion window to depict STN borders a level of 820 and a width range

of 1,080–1,180 was chosen for T2-weighted images. For SWI, a

level of 280 and a width of 264 were chosen for all images.

The parameters for T2-weighted images were as follows: rep-

etition time 5,000 ms; echo time 70 ms; slice thickness 2 mm; vox-

el size 0.7 × 0.7 × 2.0 mm; and acquisition time 8:47 min. For SWI,

the parameters were as follows: repetition time 30 ms; echo time

20 ms; slice thickness 1 mm; voxel size 0.5 × 0.5 × 1.0 mm; and ac-

quisition time 11:40 min.

Evaluation of the Stimulation Hotspot

Stereotactic coordinates of the center of the active contact at final

stimulation programming were retrieved using the co-registered iCT

used for DBS lead accuracy evaluation and defined as the center

of stimulation. The stereotactic coordinates of the center of stimula-

tion relative to the medial STN border were averaged and compared

with the predefined theoretic stimulation “hotspot” (2.6 lateral, 0.7

anterior, and 1.9 superior to the medial STN border).

Statistical Analysis

After testing for normality using the Shapiro-Wilk test (W sta-

tistic >0.900), numeric data were presented as mean with standard

deviation. We compared mean coordinates between T2 and SWI

sequences using the paired-samples T test. To compare mean co-

ordinates between the current study and the study of Bot et al. [12],

and between left and right hemisphere, we used the independent-

samples T test, after conversion of left (negative) X-coordinates

into absolute values. The average active contact coordinates were

compared to the predefined theoretic stimulation “hotspot” using

the 1-sample T test. SPSS Statistics 25 (IBM Corp., Armonk, NY,

USA) was used for all statistical analyses. A p value of <0.05 was

considered statistically significant.

Results

116 STN DBS leads were placed in 65 patients. Fifty-

one patients underwent bilateral STN DBS. Fourteen pro-

cedures were unilateral, whereof 8 in the left hemisphere

and 6 in the right hemisphere. An overview of baseline

patient characteristics is presented in Table 1. Fifty-four

patients out of 65 (83%) were included in current analysis,

providing 108 stereotactic coordinates of medial STN bor-

ders on both T2-weighted imaging and SWI. A flowchart is

presented in Figure 2, including reasons for exclusion.

![Fig. 1. Defining the medial STN border (red arrow) by drawing a line perpendicular to the AC-PC line at the anterior border of the RN; at its biggest diameter on axial plane on 3-Tesla SWI. STN, subthalamic nucleus; AC, anterior commissure; PC, posterior commissure; RN, red nucleus; SWI, susceptibility-weighted imaging.](image-url)
Table 1. Patient characteristics

Baseline clinical characteristics (n = 65)

| Characteristic                        | Value         |
|---------------------------------------|---------------|
| Female                                | 14 (22%)      |
| Age, years                            | 64 (59–67)    |
| Disease duration, years               | 9 (6–14)      |
| L-dopa responsiveness, %             | 52±19         |
| LEDD, mg                              | 1,041 (650–1,485) |

Intraoperative characteristics (n = 65)

| Characteristic                        | Value         |
|---------------------------------------|---------------|
| Bilateral STN                         | 51 (79%)      |
| Unilateral left STN                   | 8 (12%)       |
| Unilateral right STN                  | 6 (9%)        |
| Number of MER tracks                  | 1.4±0.6       |
| “First-pass” STN length, mm           | 4.8±2.1       |

Outcome (n = 42)

| Characteristic                        | Value         |
|---------------------------------------|---------------|
| LEDD, mg                              | 300 (100–510) |
| Absolute LEDD reduction, mg            | 700 (440–1,208) |
| LEDD reduction, %                      | 76 (50–88)    |

Table 2. Medial STN border coordinates on 3-Tesla T2 and SWI sequences

| Coordinate | Average coordinates medial STN border (n = 108) | p value |
|------------|-----------------------------------------------|---------|
|            | T2                                          | SWI     |
| X          | 8.9±1.0                                      | 9.0±1.1 | ns |
| Y          | −2.6±0.7                                     | −2.3±1.0| <0.001 |
| Z          | −4.0±0.5                                     | −4.5±0.7| <0.001 |

Data presented in mean ± SD. ns, not significant; STN, subthalamic nucleus; SWI, susceptibility-weighted imaging.

T2 versus SWI Sequences

Mean stereotactic coordinates of the medial STN border as defined on T2-weighted images and SWI are presented in Table 2. A statistical significant difference in means between T2-defined and SWI-defined coordinates was found in the Y-(anteroposterior) and Z-(dorsoventral) directions (paired samples t[107] = −4.1, p < 0.001 and t[107] = 7.8, p < 0.001, respectively). The anatomical location of the STN across individuals varied, as coordinates ranged from 6.1 to 10.9 mm in the X-(mediolateral) direction, from −4.3 to −1.0 mm in the Y-(anteroposterior) direction, and from −5.4 to −2.2 mm in the Z-(dorsoventral) direction on T2-weighted imaging. When defining the medial STN border on SWI, these ranges were 6.2–11.6 mm in the X-(mediolateral) direction, −4.9 to −0.1 mm in the Y-(anteroposterior) direction, and −6.0 to −2.2 mm in the Z-(dorsoventral) direction.

Left versus Right Mesencephalic Area

When defining the medial STN border on T2-weighted imaging, mean stereotactic coordinates in the Y-(anteroposterior) direction differed significantly between left and right mesencephalic area (independent t[106] = 3.6, p < 0.001; Table 3), reflecting intranidividual anatomical variability. In 44 cases (81.5%), the medial STN border was defined more anteriorly on the right side than on the left; in 10 cases (22.7%), the difference was >1.0 mm. Of note, these findings were due to the anterior border of the RN being located more anteriorly in the right mesencephalic area than in the left.

Consistently, this relative anterior location of the medial STN border in the right mesencephalic area was also found on SWI (independent t[106] = −2.3, p = 0.021; Table 4). In 10 out of the 39 patients (25.6%) with a more anterior defined medial STN border on the right mesencephalic area compared to the left, the difference was >1.0

Fig. 2. Flowchart. PD, Parkinson’s disease; STN, subthalamic nucleus; DBS, deep brain stimulation; SWI, susceptibility-weighted imaging.

Medial STN Border Reproducibility

No statistical difference in mean stereotactic coordinates of the medial STN border on T2-weighted images between the current study and the study of Bot et al. [12] was found in all 3 directions (independent t[171] = 1.6, p = 0.12 in the X-[lateral] direction; t[171] = 1.12, p = 0.26 in the Y-[anteroposterior] direction, and t[171] = 1.75, p = 0.08 in the Z-[dorsoventral] direction).

65 PD patients underwent STN DBS between October 2016 and August 2018.

Reason of exclusion:
- No SWI available (n = 3)
- History of intracranial surgery (n = 8)

64 patients included in current analysis.

Medial STN borders on T2 (n = 108)
Medial STN borders on SWI (n = 108)

Table 3. Patient characteristics

Baseline clinical characteristics (n = 65)

| Characteristic                        | Value         |
|---------------------------------------|---------------|
| Female                                | 14 (22%)      |
| Age, years                            | 64 (59–67)    |
| Disease duration, years               | 9 (6–14)      |
| L-dopa responsiveness, %             | 52±19         |
| LEDD, mg                              | 1,041 (650–1,485) |

Intraoperative characteristics (n = 65)

| Characteristic                        | Value         |
|---------------------------------------|---------------|
| Bilateral STN                         | 51 (79%)      |
| Unilateral left STN                   | 8 (12%)       |
| Unilateral right STN                  | 6 (9%)        |
| Number of MER tracks                  | 1.4±0.6       |
| “First-pass” STN length, mm           | 4.8±2.1       |

Outcome (n = 42)

| Characteristic                        | Value         |
|---------------------------------------|---------------|
| LEDD, mg                              | 300 (100–510) |
| Absolute LEDD reduction, mg            | 700 (440–1,208) |
| LEDD reduction, %                      | 76 (50–88)    |
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**Table 3. Medial STN border coordinates per brain side on 3-Tesla T2 sequences**

| Coordinate | Average coordinates medial STN border on T2 | p value |
|------------|---------------------------------------------|---------|
|            | left brain side (n = 54) | right brain side (n = 54) |         |
| X          | −9.0±1.0 | 8.9±0.9 | ns |
| Y          | −2.9±0.7 | −2.4±0.7 | <0.001 |
| Z          | −4.0±0.5 | −3.9±0.5 | ns |

Data presented in mean ± SD. ns, not significant; STN, subthalamic nucleus.

**Table 4. Medial STN border coordinates per brain side on 3 Tesla SWI**

| Coordinate | Average coordinates medial STN border on SWI | p value |
|------------|---------------------------------------------|---------|
|            | left brain side (n = 54) | right brain side (n = 54) |         |
| X          | −9.0±1.1 | 9.0±1.0 | ns |
| Y          | −2.6±1.0 | −2.1±0.9 | 0.021 |
| Z          | −4.5±0.7 | −4.4±0.7 | ns |

Data presented in mean ± SD. ns, not significant; STN, subthalamic nucleus; SWI, susceptibility-weighted imaging.

**Table 5. Stimulation “hotspot” evaluation**

| Coordinate | Average center of stimulation coordinates relative to the medial STN border | p value |
|------------|--------------------------------------------------------------------------|---------|
|            | current cohort (n = 66) | predefined “hotspot” |         |
| X          | 3.1±1.2 | 2.6 | 0.003 |
| Y          | 0.7±1.1 | 0.7 | ns |
| Z          | 1.8±1.4 | 1.9 | ns |

Data presented in mean ± SD. ns, not significant; STN, subthalamic nucleus.

**Table 6. BenGun channel used for DBS lead placement**

| BenGun channel used for DBS lead (n = 66) |  |
|------------------------------------------|---------|
| Central | 32 (48.5%) |
| Anterior | 0 (0%) |
| Lateral | 3 (4.5%) |
| Posterior | 6 (9.1%) |
| Medial | 5 (7.6%) |
| Anterolateral | 0 (0%) |
| Anteromedial | 3 (4.5%) |
| Posterolateral | 4 (6.1%) |
| Posteromedial | 4 (6.1%) |
| Stage move | 7 (10.6%) |
| Missing data | 2 (3%) |

DBS, deep brain stimulation.

mm. Taken T2 and SWI sequences together, 3 cases had a >1.0 mm anterior medial STN border on the right side compared to the left on both sequences.

**Evaluation of the Stimulation Hotspot**

Thirty-six out of the 54 included patients (66.7%) had finished stimulation programming. Sixty-six DBS leads were placed in these patients (30 bilateral, 6 unilateral), of which a majority of 41 leads (62.1%) were activated in monopolar configuration. Of the other DBS leads, 9 (13.6%) were activated in bipolar, 8 (12.1%) in double monopolar, and 8 (12.1%) in interleaving configurations. The average stereotactic coordinates of the center of stimulation relative to the medial STN border on T2 sequences were 3.1 ± 1.2 mm lateral, 0.7 ± 1.1 mm anterior, and 1.8 ± 1.4 mm superior, in proximity of the predefined theoretic stimulation hotspot (Table 5). Thirty-two (48.5%) of the DBS leads were placed using the central channel of the BenGun. Other DBS leads were placed in the posterior (n = 6), medial (n = 5), and lateral (n = 3) channel using the “+” configuration of the BenGun, and in the posteromedial (n = 4), posterolateral (n = 4), and anteromedial (n = 3) channel using the “x” configuration of the BenGun [25]. In 7 instances, adjustment of the head stage (stage move) was performed prior to DBS lead placement, since adjustment could not be achieved by using 1 of the predefined channels of the BenGun (see Table 6).

**Discussion**

In the current study, we evaluated the applicability of a new, individual anatomical reference point in STN DBS for PD: the medial STN border. Using recently described methods [12], we were able to reproduce similar stereotactic coordinates of the medial STN border in patients with PD that underwent surgery in a different center. Furthermore, the active contacts of the patients in the current study were located in close proximity of the predefined
theoretic stimulation “hotspot.” Therefore, our results suggest that the medial STN border as an anatomical reference point is suitable in order to account for individual anatomical variability in STN DBS, if one is aware of the discrepancies between T2-weighted imaging and SWI. Interestingly, our results imply that within PD patients, an anatomical difference between left and right position of the RN exists.

The Medial STN Border

In concordance with previous results [12], individual stereotactic coordinates of the defined medial STN borders varied considerably in all 3 directions, underlining the need to account for anatomical variability between patients. Above that, in the current study we found a statistical difference in mean Y-(anteroposterior) coordinates between left and right medial STN border, suggesting that accounting for anatomical variability within patients is also necessitated.

García-García et al. [11] created an adaptable 3D atlas in order to account for individual anatomical variability in determining the optimal stimulation location. However, this technique is based on the Morel atlas which does not provide bilateral anatomical information and required various software packages that may not be available for all DBS centers. Using the medial STN border as an anatomical reference point, which is easily defined by the coinciding point of the well-known Bejiani line with the delineation of the medial STN border visualized on conventional imaging, can overcome both these issues.

T2 versus SWI

Although SWI has gained interest in DBS surgery for PD for its better visualization of the STN compared to T2-weighted imaging, the reliability of SWI has been a point of discussion [6, 7, 15, 16]. For instance, discrepancies are found in defining dorsal and ventral STN borders between SWI and MER [7], especially when using lower field strengths [6]. In our current study we found discrepancies between 3-Tesla T2 and SWI sequences regarding medial STN border coordinates, that reached statistical significance in the Y-(anteroposterior) and Z-(dorsoventral) directions. Whether the use of ultrahigh field strength (7-Tesla) MRI improves the discrepancies between T2-weighted imaging and SWI remains uncertain. Therefore, when implementing the medial STN border as a new anatomical reference in STN DBS, 1 should be aware of discrepancies in visualization of the basal ganglia between T2-weighted imaging and SWI, for instance in defining an optimal stimulation location or localizing kinesthetic cell clusters. Since the electrophysiological medial STN border is generally not sought using MER in DBS surgery for PD, differences in reliability between T2-weighted imaging and SWI could not be measured in the current study.

The Stimulation Hotspot

Although a statistical difference in means was found in the X-(lateral) direction between the theoretic stimulation “hotspot” defined by Bot et al. [12] and the stimulation location of the patients in the current study, we consider this negligible for 2 reasons. First, the coordinates represent a center within an electric potential field rather than a pinpoint “hotspot.” The volume of tissue activated likely exceeds 0.5 mm and, above that, the diameter of a contact is more than 1 mm. Second, both stimulation locations were found in the dorsolateral part of the STN. Taken together, this suggests that the stimulation “hotspot” (located 2.6 mm lateral, 0.7 mm anterior, and 1.9 mm superior to the medial STN border) is applicable in PD patients that underwent STN DBS surgery across centers.

Alternatively, the stimulation “hotspot” could be evaluated by correlating the 3-dimensional distance between the hotspot coordinates and the center of stimulation of the patients with an UPDRS-III reduction at follow-up. At the senior authors’ institution, however, the UPDRS-III is not standardized at follow-up, given the vast majority of the patients have to travel a considerable amount of time and assessment of the UPDRS-III is time consuming. Therefore, the LEDD reduction at follow-up is used to evaluate the effectiveness of stimulation. However, the LEDD reduction is not a lateralized outcome, whereas the majority of PD patients that undergo DBS surgery receive bilateral treatment. In case of bilateral stimulation, it is not possible to determine to what extent stimulation in each hemisphere contributes to reduction in LEDD. We, therefore, considered the LEDD reduction at follow-up not suitable as a surrogate outcome measure. The reduction in lateralized UPDRS-III score seems suitable as an outcome measure, although stimulation in 1 hemisphere can have clinical effect in the ipsilateral body side as well [26, 27].

Andrade-Souza et al. [28] concluded that targeting based on the borders of the RN is more accurate compared to the established indirect targeting (AC-PC based) and modified direct targeting (3 mm above the center of the STN on coronal sections using T2-weighted images). As the authors point out, the distance between their RN-
based target and the location of the optimal contact was nonetheless more than 3 mm on average and therefore electrophysiological determination of the final electrode position was recommended. Our patient-specific reference point (also RN based) allows to further optimize targeting within the dorsolateral STN. We, therefore, encourage other centers to localize a stimulation “hotspot” by calculating the stereotactic distance between active DBS lead contacts and the medial STN border using optimal responders within their patient population, and compare the results to the predefined theoretic stimulation “hotspot” by Bot et al. [12] (2.6 mm lateral, 0.7 mm anterior, and 1.9 mm superior to the medial STN border on T2-weighted imaging). Eventually, this could enable direct medial STN border-based targeting using data from optimal responders worldwide and increase overall clinical effectiveness of STN-DBS in future patients with Parkinson’s disease.

The Red Nucleus

Since we defined the medial STN border using the anterior border of the RN at its maximal diameter, our findings of a statistically different average left and right Y-coordinate, on both T2-weighted imaging and SWI, directly reflect a more anterior position of the anterior border of the RN in the right mesencephalic area compared to the left. In roughly one out of 4 patients, the right RN was more than 1 mm anterior than the left, which we consider clinically relevant in stereotactic DBS surgery.

These findings raise the question to what extent the anatomical position of the RN (mesencephalic origin) and the STN (diencephalic origin) relate. Also, it remains uncertain whether these findings are disease and/or age related. All the patients included in current study were right handed (data not provided).

One study of Xiao et al. [14] investigated morphometric variability of the STN and RN in patients with advanced PD. In concordance with our findings, the authors report a more anteriorly mean distance between the geometric center of the right RN to the MCP than the left RN. Similar differences were found regarding the STN, also when using the center of mass to calculate the distance to the MCP; however, left and right measurements were not directly statistically compared to each other but independently to the Talairach [29] atlas.

Various reports provide data on left and right volumetric properties of basal ganglia, including the STN and RN [10, 14, 30–32]. However, analysis of volumetric variability between left and right hemispheres was either not significant [10, 30, 31] or did not lay within the scope of the study [32]. In theory, however, a volumetric change could occur independent from positional or morphometric changes.

Study Limitations

Although the use of intraoperative CT to determine lead placement accuracy in deep brain stimulation surgery has been validated [33–35], inherent measurement error has been reported [34]. Therefore, fusion of preoperative MRI and iCT images were carefully inspected for each case to reduce merging error. In addition, standard surgical precautions were taken to limit the effect of brain shift on lead placement accuracy [36, 37]. Finally, it should be noted that the difference in slice thickness values of our study potentially limits our methodology. It has previously been suggested, however, that these values would provide a comparable degree of accuracy [7, 38].

Conclusion

The medial STN border is applicable across centers as a new anatomical reference point for STN DBS surgery for PD, and seems suitable in order to account for interindividual and intrindividual anatomical variability if one is aware of the discrepancies between T2-weighted imaging and SWI. The clinical implications of our findings of a more anteriorly positioned RN in the right mesencephalic area compared to the left should be further investigated.

Statement of Ethics

The study was approved by the Rush University Institutional Review Board (ID: 18050108; date of approval: June 11, 2018), and informed consent was obtained.

Conflict of Interest Statement

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Author Contributions

E.B.: conceptualization, methodology, investigation, data curation, formal analysis, visualization, and writing – original draft. M.B.: conceptualization, methodology, and writing – review and editing. P.V.D.M.: project administration, conceptualization, methodology, and writing – review and editing. G.P.: conceptualization, data curation, and writing – review and editing. S.S.: software, resources, data curation, and writing – review and editing. L.V.M.: project administration, conceptualization, methodology, data curation, supervision, and writing – review and editing.

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