Current Directions in Deep Brain Stimulation for Parkinson’s Disease—Directing Current to Maximize Clinical Benefit

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ABSTRACT

Several single-center studies and one large multicenter clinical trial demonstrated that directional deep brain stimulation (DBS) could optimize the volume of tissue activated (VTA) based on the individual placement of the lead in relation to the target. The ability to generate axially asymmetric fields of stimulation translates into a broader therapeutic window (TW) compared to conventional DBS. However, changing the shape and surface of stimulating electrodes (directional segmented vs. conventional ring-shaped) also demands a revision of the programming strategies employed for DBS programming. Model-based approaches have been used to predict the shape of the VTA, which can be visualized on standardized neuroimaging atlases or individual magnetic resonance imaging. While potentially useful for optimizing clinical care, these systems remain limited by factors such as patient-specific anatomical variability, postsurgical lead migrations, and inability to account for individual contact impedances and orientation of the systems of fibers surrounding the electrode. Alternative programming tools based on the functional assessment of stimulation-induced clinical benefits and side effects allow one to collect and analyze data from each electrode of the DBS system and provide an action plan of ranked alternatives for therapeutic settings based on the selection of optimal directional...
contacts. Overall, an increasing amount of data supports the use of directional DBS. It is conceivable that the use of directionality may reduce the need for complex programming paradigms such as bipolar configurations, frequency or pulse width modulation, or interleaving. At a minimum, stimulation through directional electrodes can be considered as another tool to improve the benefit/side effect ratio. At a maximum, directionality may become the preferred way to program because of its larger TW and lower energy consumption.

Keywords: Contact; Deep brain stimulation; Directionality; Lead; Parkinson disease; Programming

Key Summary Points

Directional deep brain stimulation (DBS) has the potential to minimize stimulation-induced side effects and maximize clinical benefits.

To maximize the opportunity of directionality, the surgical planning should be based on the DBS lead level containing segmented directional electrodes.

Visualization software platforms can assist programming by estimating the volume of tissue activated by conventional or directional DBS electrodes.

Functional software platforms can support directional programming by creating an action plan of ranked alternatives that may be needed over time.

Directional DBS may avoid the need for complex stimulation protocols, such as bipolar stimulation, frequency or pulse width modulation, or interleaving.

INTRODUCTION

Over the last three decades, the long-term efficacy of deep brain stimulation (DBS) in the management of Parkinson disease (PD) has been well established, with 60% improvement of levodopa-related motor complications [1, 2], 40–60% amelioration in quality of life [3], and a significant increase in quality-adjusted life expectancy [4]. Despite these remarkable results, not all patients experience maximal benefit from DBS. Unintentional spread of stimulation outside the target and into adjacent areas may result in dose-limiting motor or non-motor side effects, preventing an optimal outcome [5, 6].
Directional leads represent an innovative technology designed to address this issue. The concept of steering current away from side effects and towards benefits after the lead is locked into place is appealing when considering the need to further improve the accuracy of DBS targeting [7]. Here, we discuss opportunities and challenges for directional DBS and review data in support of the integration of directional leads into clinical practice with regards to the functional anatomy of DBS target structures, planning strategies for optimal lead placement, and use of software platforms to assist the selection of the most promising stimulation settings. This article is based on previously conducted studies and does not contain any studies with human participants or animals performed by any of the authors.

**CLINICAL DATA SUPPORTING DIRECTIONALITY**

In 2014, two groups reported the results of intraoperative studies of directional DBS leads (not permanently implanted) (Table 1). In one study [8], the therapeutic window (TW), defined as the range between the minimal intensity of stimulation required to obtain meaningful clinical benefits, or therapeutic current strength (TCS), and the intensity of stimulation at which the first persistent side effect occurred, or side effect threshold (SET), was evaluated for single directional and omnidirectional configurations in 13 patients with PD or essential tremor (ET). The electrode used had two segmented contacts at the distal end, with three segments each, and two solid ring electrodes proximally. When test stimulation was performed, the TW was found to be 41.3% wider using the best directional electrode, compared to omnidirectional stimulation at that same level. The TCS was 43% lower for the best directional electrode compared to that required for omnidirectional stimulation, indicating a higher degree of efficiency with directional DBS. In the second study [9], eight patients with PD were assessed intraoperatively using an experimental lead with 32 oval disc-shaped thin-film electrode contacts arranged on eight rows of four contacts. The patient, the evaluating neurologist, and the neurosurgeon were blind to the direction of the steering modes. When compared to omnidirectional stimulation, directional DBS increased the TW in all patients, with a gain ranging from 0.5 to 1.5 mA. In seven out of eight cases, SETs could be increased by at least 1 mA when using directional DBS.

Three single-center studies investigated US Food and Drug Administration (FDA)-approved directional implants (Table 1) employing a “1-3-3-1” lead design with the two central rows segmented into three individually controlled electrodes; each segment spaced 120° apart in the plane orthogonal to the long axis of the lead. These studies found that best directional DBS has a significantly larger median TW and higher SET [10, 11], as well as slightly lower total electrical energy delivered (TEED) [12]. More recently, a large international, multicenter, prospective, crossover study of 234 patients with PD undergoing subthalamic nucleus (STN) DBS from 37 centers across seven countries (the PROGRESS Study, Abbott St. Jude Medical, Plano, TX, USA) confirmed the superiority of directional vs. omnidirectional programming [13]. In this study, patients were conventionally (i.e., non-directionally) stimulated for 3 months at the optimal ring. At the 3-month primary endpoint, the optimal ring settings were confirmed and in the same session the best directional settings were identified between the three individual segments and the three pairwise combinations of segments at the best segmented level. The best ring and best directional montages were then evaluated in a randomized (based on a coin toss) and double-blinded (patient and the clinician) fashion and their TWs were recorded. Study details and results of the primary endpoint are included in Table 2. All patients had 12 months of follow-up and several measures were tracked in this period at 3, 6, and 12 months including Unified PD Rating Scale (UPDRS)-III (motor section), UPDRS-II (activities of daily living section), clinician preference, patient preference, 39-item PD Questionnaire (PDQ-39), adverse events, and others.
| Author and year | Sample size | Lead configuration | Clinical setting | Efficacy measures | FUP | Main results |
|-----------------|-------------|--------------------|------------------|------------------|-----|--------------|
| Pollo et al. 2014 [8] | 11 PD (STN) 2 ET (Vim) 2 distal segmented contacts (3 segments each) 2 proximal ring contacts | 2E T (Vim) 2 proximal ring contacts | Intraoperative double-blinded evaluation | Full effect on rigidity or good effect on tremor | NA | TW 41.3% wider and TCS 43% lower with directional vs. omnidirectional stimulation |
| Contarino et al. 2014 [9] | 8 PD (STN) 32 oval disc-shaped contacts | | Intraoperative double-blinded evaluation | Full effect on rigidity | NA | TW wider (0.5–1.5 mA) with directional vs. omnidirectional stimulation |
| Steigerwald et al. 2016 [10] | 7 PD (STN) 2 central segmented contacts (3 segments each) 1 proximal and 1 distal ring contacts | | Retrospective unblinded analysis of permanently implanted patients | Full effect on rigidity | 3–6 months | TW variations from −100% to +440% with directional vs. omnidirectional stimulation Best TW improvement with the best directional contact at the less effective level At 3–6 months, all patients remained in directional stimulation |
Table 1 continued

| Author and year | Sample size | Lead configuration | Clinical setting | Efficacy measures | FUP | Main results |
|-----------------|-------------|--------------------|------------------|------------------|-----|--------------|
| Dembek et al.   | 10 PD (STN) | 2 central segmented contacts (3 segments each) 1 proximal and 1 distal ring contacts | Prospective double-blinded | Improvement of at least 1.5 points in the UPDRS-III composite scores of upper limb rigidity, finger tapping, and hand rotation | 3–6 months | TW wider with directional vs. omnidirectional stimulation (median 2 mA vs 1 mA) SET higher with directional vs. omnidirectional stimulation (median 4 mA vs 3 mA) At 3–6 months, 14 leads remained in directional stimulation |
| Rebelo et al.   | 3 PD (Vim)  | 2 central segmented contacts (3 segments each) 1 proximal and 1 distal ring contacts | Retrospective unblinded analysis of permanently implanted patients | Full effect on tremor | 6 months | TW wider (1.86 mA vs. 0.97 mA) and TCS lower (1.51 mA vs. 2.19 mA) with directional vs. omnidirectional stimulation TEED 6–18% lower with directional vs. omnidirectional stimulation At 6 months, 9 leads remained in directional stimulation |

*DT* dystonic tremor, *ET* essential tremor, *FUP* follow-up, *PD* Parkinson disease, *NA* not applicable, *SET* side effects threshold, *STN* subthalamic nucleus, *TCS* therapeutic current strength, *TEED* total electrical energy delivered, *TW* therapeutic window, *UPDRS* Unified PD Rating Scale, *Vim* ventral intermediate nucleus
CHALLENGES ASSOCIATED WITH DIRECTIONAL PROGRAMMING

While directional DBS systems allow fine control over the parameters by which they deliver stimulation, including location, size, shape, and nature of neural activation, this increased ability to provide accurate and personalized stimulation demands a revision of paradigms currently used for the placement and programming of DBS electrodes. In particular, the anatomical and functional connections of the DBS targets, as well as the bioelectrical properties of directional vs. conventional leads, represent critical aspects to be considered for the development of innovative DBS protocols.

DBS Target Structures: Anatomical and Functional Connections

The STN is a small lens-shaped structure located in the anterior-lateral portion of the midbrain at the junction of the cerebral peduncle to the tegmentum (Fig. 1). Leads are generally implanted following an oblique trajectory from

| Table 2 | Data from the PROGRESS study |
|---------|-----------------------------|
| Device  | Abbott St. Jude Infinity™ IPG system |
| Enrollment | 234 patients with PD (157 male; 77 female) |
| Demographics | Age 61.7 ± 8.4 years |
|            | PD duration since onset 11.7 ± 7.6 years |
|            | PD duration since diagnosis 10.2 ± 7.4 years |
| Number of centers | 37 centers, from 7 countries |
| Lead configuration | 2 central segmented contacts (3 segments each), 1 proximal and 1 distal ring contacts; 1-3-3-1 |
| Clinical setting | Prospective, blinded subject, blinded observer, crossover study of directional versus non-directional stimulation |
| Study endpoints | Primary endpoint |
|            | Superiority benchmark: at least 60% of patients at 3 months have wider TW with directional stimulation when compared to non-directional stimulation in a randomized evaluation |
|            | Secondary endpoints |
|            | Non-inferiority benchmark: 40–60% of patients at 3 months have wider TW with directional stimulation when compared to non-directional stimulation in a randomized evaluation |
|            | Comparison of UPDRS III at 3 months on non-directional stimulation versus at 6 months after switching to directional stimulation |
|            | Descriptive endpoints |
|            | Patient and clinician preference at 6 months between directional and non-directional stimulation |
|            | Comparison of TW and TCS amplitudes between directional and non-directional stimulation |
|            | Device-related adverse events |
| Primary endpoint results | In 90.6% of patients (183/202), TW was wider with directional stimulation as compared to non-directional stimulation; primary endpoint superiority exceeded ($p < 0.001$) |

DBS deep brain stimulation, PD Parkinson disease, TCS therapeutic current strength, TW therapeutic window

△ Adis
a dorsal lateral entry point to a more ventral medial endpoint in the STN. Given the high number of important surrounding structures, stimulation-induced side effects are important to recognize and, if seen, can guide inferences about the location of the lead in relation to the planned STN target (Table 3). Functionally, the STN has a complex microanatomy, with large dendritic trees that are synaptically convergent between afferents from different functional areas [14–16] (Table 4).

The globus pallidus pars interna (GPI) is a triangular-shaped structure forming the inner part of the lentiform nucleus of the basal ganglia (Fig. 1). DBS leads are generally implanted following a parasagittal trajectory that traverses putamen, globus pallidus pars externa (GPe), and GPI. The target is in the posterolateral part of GPI, just medial to the medullary lamina that separates GPe and GPI and the tip of the electrode will be just dorsolateral to the optic tract. Stimulation-induced side effects, when present, can guide inferences about the location of the lead in relation to the planned target (Table 3). Functionally, the GPI is a site of convergence of striatal–pallidal efferents [17–28]. Together with the substantia nigra pars reticulata (SNr), GPI represents the main basal ganglia output to the ventroanterior and ventrolateral thalamic nuclei and, eventually, to the cortex [29] (Table 4).

Surgical Approach and Considerations Related to Type of DBS Lead

In currently used planning strategies, the STN may be targeted at the anterior border of the red nucleus, 2 mm lateral to its medial border, and at 4 mm inferior to the anterior commissure–posterior commissure (AC–PC) plane [30]. GPI may be targeted 3 mm lateral to its medial border and one-third of its length anterior from its posterior margin, all at the AC–PC plane [30]. However, these heuristics are not absolute, and other effective targeting strategies may also be applied. In addition, the precise targets within these areas are still contested by experts [31, 32]. In particular, the optimal spot for stimulation within the STN remains debated [33].

For directional leads, targeting currently remains the same as for legacy quadrripolar leads. However, more attention must be paid to the depth of the electrodes. Most manufacturers (Boston Scientific and Abbott) have directional capabilities only in their second and third contact rows, not in the first or fourth. The Aleva directSTN Acute lead, on the other hand, has directional first and second rows only, which better suit GPI because of the proximity to the internal capsule at the bottom of the lead.

Fig. 1 DBS target structures. Anatomical DBS structures of the left hemisphere in the anterior (left) and posterior (right) views. Courtesy of Abbott’s anatomical visualization educational software (StimDirect), available on the St. Jude Infinity™ clinician programmer. Subthalamic nucleus (STN): ventrally, the STN is bordered by the substantia nigra (SN), anterolaterally by the internal capsule (IC), posteriorly by the medial lemniscus (ML), dorsally by the zona incerta and the fields of Forel, and medially by the red nucleus (RN), the medial forebrain bundle, and the midbrain course of the oculomotor nerve. Globus pallidus pars interna (GPI): ventrally, the GPI is bordered by the ansa lenticularis, which separates it from the nucleus basalis and the amygdala, ventromedially by the optic tract, dorsally and medially by the posterior limb of the internal capsule, and laterally by the internal medullary lamina of the globus pallidus which separates it from the globus pallidus pars externa (GPe). Additionally, the GPI is divided into an internal and external component by the incomplete medullary lamina of the globus pallidus. Thalamus

△ Adis
With currently available electrodes, the depth of the electrode should be targeted with the intention of using a directional row (typically the second or third) as the active contact. Depending on the surgeon’s practice, this might require inserting the electrode deeper than with a standard quadripolar lead.

To help ensure consistent orientation of the segments of the directional lead, manufacturers have placed a fiducial on the lead proximal to the four electrode levels. When the fiducial points in the anterior direction, the three segments face anteriorly, posteromedially, and posterolaterally, respectively. Nevertheless, during implantation, the lead is subject to torsional movements, making it difficult to predict the exact position of the fiducial marker and, therefore, the exact orientation of each segment. If there is uncertainty regarding the directionality, high-resolution X-rays, fluoroscopy, or CT scan can be used to verify the rotational orientation [35].

### DIRECTIONAL PROGRAMMING

The main goal of the first programming visit is to determine the TW for each of the electrode segments. This process involves

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Table 3  Stimulation-induced side effects

| Direction of current spread | Side effect | Structures involved |
|----------------------------|------------|---------------------|
| STN                        | Lateral or anterolateral | Contralateral muscle contractions | Internal capsule |
|                            |            | Facial or tongue pulling | |
|                            |            | Dysarthria | |
|                            |            | Eyelid opening apraxia | |
|                            |            | Contralateral gaze deviation | |
| Anteromedial               |            | Autonomic changes/vegetative side effects | Lateral hypothalamic area |
|                            |            | (nausea, heat sensation, sweating) | |
| Medioventral               |            | Disconjugate gaze | Oculomotor nerve |
| Posterior                  |            | Paresthesias | Medial lemniscus |
| Dorsal or ventral          |            | Bradykinesia worsening | Zona incerta/thalamus |
|                            |            | Levodopa effect reduction | Substantia nigra |
|                            |            | Mood changes (mania, depression, or apathy) | |
| Scarcely localizable       |            | Impulsivity | – |
|                            |            | Hypophonia | |
| GPi                        | Medial     | Contralateral muscle contractions | Internal capsule |
| Posterior                  |            | Bradykinesia worsening | |
| Dorsal                     |            | Dyskinesia | Globus pallidus pars externa |
| Ventral                    |            | Phosphenes | Optic tract |
| Lateral                    |            | No side effects | Globus pallidus pars externa |
| Anterior                   |            | | |

GPi globus pallidus pars interna, STN subthalamic nucleus

* In spite of rigidity improvement

[34]. With currently available electrodes, the depth of the electrode should be targeted with the intention of using a directional row (typically the second or third) as the active contact. Depending on the surgeon’s practice, this might require inserting the electrode deeper than with a standard quadripolar lead.

To help ensure consistent orientation of the segments of the directional lead, manufacturers have placed a fiducial on the lead proximal to the four electrode levels. When the fiducial points in the anterior direction, the three segments face anteriorly, posteromedially, and posterolaterally, respectively. Nevertheless, during implantation, the lead is subject to torsional movements, making it difficult to predict the exact position of the fiducial marker and, therefore, the exact orientation of each segment. If there is uncertainty regarding the directionality, high-resolution X-rays, fluoroscopy, or CT scan can be used to verify the rotational orientation [35].
Contacts [36]. Directional leads are often first tested in ring mode, followed by directional stimulation on individual segments if efficacy in ring mode is limited by stimulation-related side effects, or to identify programming strategies that allow a larger TW with lower energy consumption [37]. Supporting programming software allows for visualizations of the anatomical context along with the predicted volume of tissue activated (VTA) and for functional mapping of the thresholds for clinical efficacy and side effects. Some of these tools also allow for an integrated analysis of power consumption adjusted to each contact impedance.

### Table 4  STN and GPi functional connections

| Input  | Neurotransmitter | STN area involved | Output  | Neurotransmitter |
|--------|------------------|--------------------|---------|------------------|
| STN    | GABA             | Dorsolateral (motor area) | Globus pallidus pars interna | Glutamate |
|        |                  | Ventromedial (limbic area) |          |                  |
| Primary motor cortex | Glutamate | Dorsolateral (motor area) | Substantia nigra pars reticulata | Glutamate |
| Supplementary motor area |          | Ventromedial (limbic area) |          |                  |
| Premotor cortex |          |          |          |                  |
| Prefrontal cortex | Glutamate | Ventromedial (limbic area) |          |                  |
| Prelimbic-medial orbital areas |          |          |          |                  |
| Thalamus (parafascicular and centromedian nuclei) | Glutamate | Dorsolateral (motor area) |          |                  |
|        |                  | Ventromedial (limbic area) |          |                  |

### Table 4  STN and GPi functional connections (continued)

| Input  | Neurotransmitter | STN area involved | Output  | Neurotransmitter |
|--------|------------------|--------------------|---------|------------------|
| GPi    | GABA             | Thalamus (ventroanterior and ventrolateral nuclei) | GABA   |
|        |                  |                    |         |                  |
| Globus pallidus pars externa | GABA | Substantia nigra pars compacta | GABA   |
| Thalamus (intralaminar nucleus) | Glutamate | Lateral habenular nucleus | GABA   |
| STN    | Glutamate        | Pedunculopontine nucleus | GABA   |
| Pedunculopontine nucleus | Glutamate and acetylcholine | – | – |
| Dorsal raphe nucleus | Serotonin | – | – |
| Substantia nigra | Dopamine | – | – |

GABA gamma-aminobutyric acid, GPi globus pallidus pars interna, STN subthalamic nucleus
Visualization of Volume of Tissue Activated

The volume of neural activation can be estimated using finite element techniques, wherein a computer system subdivides a volumetric region of interest and solves for a scenario by applying appropriate mathematical relationships, initial conditions, and boundary conditions and iterating until a stable solution results. In this domain, this is applied by positioning an appropriate model of a stimulating electrode, configuring it with appropriate stimulation parameters (polarities, stimulus amplitude), and determining the resulting current flows and voltage gradients in the surrounding tissue. These electromagnetic gradients can then be applied, in a second step, to a population of simulated neurons positioned and oriented preferentially around the stimulating electrode so as to determine points at which neurons may be activated by a given stimulation regime [38]. These neurons may be modeled such that they respond both to the intensity of the stimulation (current amplitude) and the time-dependent parameters such as the width of a stimulation pulse [38]. As an example, various neuron fiber diameters may be included, such that the response of smaller fibers to a given stimulus pulse is different than that of larger fibers. The resulting neuron population that is predicted to be activated can then be represented visually, as a volume enclosing the points of activation.

Model-based approaches have been shown to have some predictive value when used in visual programming systems [39]. However, validating these models can be challenging, as it is difficult to know with certainty which neural tissue volumes are indeed activated by stimulation. One thing that has been established is that fully featured models incorporating voltage drop and capacitance of the electrode–electrolyte interface, tissue encapsulation of the electrode, and diffusion tensor-based 3D tissue anisotropy and inhomogeneity produce more realistic predictions than simpler models that do not account for these factors [40].

As new stimulation modalities are introduced, including the new options afforded by directional leads, these VTA models may be used to compare the different stimulation options available. As examples, in bipolar stimulation, they can be used to understand the differences in volume between anodal and cathodal activation regions. For interleaved stimulation, they can be used to understand areas of overlap in separate pulse trains, wherein tissue receives stimulation at double the underlying frequency [41]. In directional stimulation, the ability to push stimulation off axis, commonly referred to as displacement of the VTA centroid, can be explored in comparison to traditional ring mode stimulation. This may be of specific interest, as the thresholds to activation in directional stimulation may be different than in traditional rings. For fixed stimulation amplitudes, the smaller surface area of a directional segment creates relatively higher gradients around the electrode which may increase the probability of neural activation [38].

Visualization Strategies: Opportunities

Patient-Specific Anatomy
A preoperative magnetic resonance imaging (MRI) may be used to establish the location and boundaries of anatomical structures, either through adaptation of an anatomical atlas or direct segmentation of structure boundaries from intensity information in the images themselves. Further, fiber tracts may additionally be represented if appropriate imaging sequences (diffusion tensor imaging) or atlases are available.

Final Lead Location and Orientation
A postoperative, or in some cases an intraoperative, image may be used to establish the final lead location and orientation. For this purpose, images that clearly delineate the electrode positions and the orientation of fiducial markers included in the lead body via a hyper- or hypo-intense image artifact can be used.

Visualization of Predicted Stimulation Extent
A VTA or similar model may be applied to the representation of the stimulating electrodes. Models may be created for one or more
potential sets of stimulating parameters (i.e., different active contacts, different current or voltage amplitudes, monopolar vs. bipolar stimulation, directional vs. conventional ring stimulation, etc.) such that the overlap of the resulting stimulation field to patient anatomy can be assessed.

Integration of Ancillary Information
In addition to anatomy, lead location, and stimulation extent, visualization solutions may additionally allow for the addition of other information of potential use in identifying optimal stimulation settings. For example, intra- or postoperative electrophysiological measures may be visualized in the patient-specific anatomical context. In addition, aggregated historical information about stimulation outcome in the form of a statistical outcome or side effect map may be visualized in the patient-specific anatomical context to further inform programming decisions.

Visualization Strategies: Challenges

Limitations of Anatomical Models
Currently used anatomical models on which the lead is visualized are usually not patient-specific, and even when patient-specific, they are based on presurgical MRI images that do not account for procedural brain shift and postsurgical anatomical changes reported at up to 4 mm in the deep brain [42–46].

Deviation of DBS Lead
DBS leads show large deviations from their intended implanted orientation: more than 30° rotation in 42% of the leads and more than 60° rotation in 11% of the leads [47]. Thus, the orientation of the individual segmented contacts might be no longer valid relative to the underlying anatomy in presurgical MRI presented during programming. Furthermore, significant lead migration (greater than 3 mm) along the ventrodorsal axis or upward displacement from immediate to delayed CT has been reported in over 12% of leads placed at an expert center [48, 49].

Variability in Tissue Impedance and Anisotropy
Validating VTA models is challenging as it is difficult to know with certainty which neural tissue volumes are stimulated [50]. VTA models made available for programming are based on generic homogeneous models that do not account for tissue inhomogeneities, for instance permittivity and conductivity of brain structures, which may alter VTA predictions from −44% to 174% [51]. Also, VTA models available in programming platforms do not account for patient-specific tissue anisotropy that can at best only be modeled using tractography from high-resolution patient-specific diffusion tensor imaging (DTI) [52, 53]. Other real-world factors, not usually modeled, that influence the accuracy, shape, and extent of the VTA include physiology and pathophysiology [50], brain pulsation and patient hydration [54], glial scar formation around the electrode [55], and local fluid retention [56].

Software Platforms to Assist DBS Programming
SureTune™ (Supplementary Fig. S1) is a programming visualization tool available for clinical use in many regions and close to being released also in the USA. It incorporates the Bardinet–Yelnick anatomic atlas and registration algorithm [57], and further allows a user to manually adjust structure size, shape, and location to allow for valid representation of a patient’s specific anatomical variation. It also supports intensity-based segmentation of MRI visible structures. Within the existing software, leads can be placed according to stereotactic coordinates or aligned via postoperative images. Anatomic representations of structures can be validated by co-registration of microelectrode recordings to confirm anatomical boundaries if additional confidence is desired. Stimulation modeling is available for common stimulation configurations, and stimulation plans can be created for use in the clinical setting. Finally, via a service offering, statistical maps of outcome or side effects can be created which can then be prospectively visualized to inform future
clinical decision-making or research applications: a recent study [58] showed that such statistical maps correlated well with best clinical programming, demonstrating predictive utility in prospective cohorts for GPi stimulation in dystonia.

While potentially useful for optimizing clinical care, SureTune™ has several limitations. Its atlas is a single brain histologically based example that may not account for subject to subject variation in anatomy. While its approach of algorithmic fitting plus human adjustment can be highly accurate, it is also dependent on image quality and human judgment to achieve a good patient-specific fit. Finally, it uses homogeneous and isotropic assumptions in its VTA methodology, which may introduce errors in the visualization due to variation of tissue properties within a specific patient’s brain.

SureTune™ is tightly integrated with other Medtronic surgical tools, allowing import of surgical information from Stealth™ navigation and planning software and the documentation of intraoperative electrophysiology which may further validate a patient-specific visualization. Medtronic’s current programming offering allows visualization of the VTA created by the device and the annotation of clinical observations. Future updates to SureTune™ will extend interoperability, allowing anatomical and electrophysiological information to appear on the programming tablet along with the stimulation volumes.

**Boston Scientific GUIDE™ XT**

The Boston Scientific neuromodulation stimulators use a system that includes both VTA modeling and functional testing. Visualization of VTAs with the Boston Scientific system is available through a software tool (GUIDE™ XT). GUIDE™ XT is compatible with the BrainLab Elements (Munich, Germany) surgical planning software, available for clinical use in many countries. The software enables 3D modeling and visualization of the VTA relative to the patient’s anatomical structures. To generate a model of lead location in the brain, a simulated DBS lead from a patient’s postoperative CT scan is registered to an anatomical atlas based on segmentation of a patient’s preoperative MRI. Clinical stimulation parameters can then be programmed onto the simulated lead to generate the associated VTA. In cases of suboptimal electrode placement or clinical response, additional options may be needed, and current steering between directional segments may be useful to expand the TW (Supplementary Fig. S2) [10]. Visualization tools may be useful for understanding these nuances of directional DBS and adjusting stimulation appropriately. For example, the VTA created by coactivation of two segments has a different shape than the VTA created by activation of a single segment, with the former having a lesser radial extent of activation than the latter at the same amplitudes. The Vercise™ system also includes the Neural Navigator software, intended to aid in functional testing of different stimulator settings by mapping the resulting clinical effects. Therapeutic effects and side effects associated with monopolar stimulation on both ring and directional electrodes are represented on a 2D clinical effects map (Supplementary Figs. S3 and S4), and effects may be recorded on all leads for all stimulation configurations. VTAs for both standard DBS leads and directional DBS leads are also visualized within the Neural Navigator software, which can import the anatomical structures from the BrainLab platform.

**Abbott Informity™**

While the Abbott Infinity™ clinician programmer does not have a VTA visualization tool, it includes the Informity™ programming software designed primarily for simplifying functional programming. The software guides the user through directional programming using visual representations of stimulation responses, occurrences of stimulation-induced symptom relief and side effects, to create an action plan of ranked electrode montage alternatives that may be needed over time during therapy. The layout and workflow of the Informity™ programming software enable “event markers” to document the amplitudes that produce symptom relief and side effects (Supplementary Figs. S5, S6, and S7). After monopolar survey of ring and segmented electrodes, the Informity™ software allows clinicians to rank the investigated
montages according to various clinically relevant criteria. With traditional omnidirectional programming, amongst montages with comparable outcomes some clinicians prioritize maximizing the TW while others prioritize minimizing the TCS (power consumption). To that end, the decision support tool available within the Informity™ software not only allows ranking the montages on the basis of power (microwatts) and TW (milliamps) but also on the basis of a measure referred to as the therapeutic window percentage (TW%). TW%, which is TW expressed as a percentage of TCS, is a means to balance the trade-off between maximizing TW and minimizing TCS and gives clinicians a means to optimize gains in both TCS and TW simultaneously. A final sorting option, balanced threshold, gives clinicians an additional level of optimization to balance gains in TW% while minimizing power consumption as per clinician preference (Supplementary Fig. S7). The software allows complete customization of symptom relief and side effect lists and enables the export of the entire functional programming session in the form of a PDF report that can be exported to electronic medical records for future review.

CONCLUSIONS

Increasing clinical evidence supports the superiority of directional over omnidirectional stimulation. It is conceivable that directional leads, by providing asymmetric stimulation, may avoid the need for complex programming paradigms such as bipolar stimulation, frequency or pulse width modulation, or interleaving. At a minimum, stimulation through directional electrodes can be considered as another tool to improve the benefit/side effect ratio. At a maximum, directional stimulation may become the preferred way to program, even when omnidirectional stimulation causes no side effects, because of its larger TW and lower energy consumption.

However, as new DBS systems increase in their capability to control the shape, size, and location of stimulated neural tissue, the complexity of optimizing the configuration of such systems grows as well. The use of simple programming protocols, such as the activation of a single directional segment, has proved to be effective in the vast majority of cases [13]. Directional multiple-electrode programming configurations, whether using independent current controllers or interleaving, may provide additional nuances in programming strategies, but may also result in increased programming complexity [59], changes to battery power consumption, and reduction in the laterality of the volume of activation [60]. To assist with DBS programming, DBS producers developed innovative software platforms for functional mapping, as well as tools for visualization of the volume of neural activation. Still, these models are constrained by simplifying assumptions, such as using homogeneous and isotropic tissue environments and arbitrarily positioned populations of generally linear axons. Also, no study has so far established if commercially available software platforms are superior to standard programming.

In conclusion, while directionality offers a unique opportunity to improve the functional outcomes of DBS, it also requires definition of unified strategies for functional mapping and common standards for VTA visualization since suboptimal programming remains one of the major causes for DBS failure [6, 61]. Here, we advocate for intensive collaboration between DBS manufacturers and academic centers to develop and test in large multicenter clinical trials innovative programming algorithms aiming at maximizing the benefits of directional DBS.

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