Complication rates, lengths of stay, and readmission rates in “awake” and “asleep” deep brain simulation

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OBJECTIVE As the number of deep brain stimulation (DBS) procedures performed under general anesthesia (“asleep” DBS) increases, it is more important to assess the rates of adverse events, inpatient lengths of stay (LOS), and 30-day readmission rates in patients undergoing these procedures compared with those in patients undergoing traditional “awake” DBS without general anesthesia.

METHODS All patients in an institutional database who had undergone awake or asleep DBS procedures performed by a single surgeon between August 2011 and August 2014 were reviewed. Adverse events, inpatient LOS, and 30-day readmissions were analyzed.

RESULTS A total of 490 electrodes were placed in 284 patients, of whom 126 (44.4%) underwent awake surgery and 158 (55.6%) underwent asleep surgery. The most frequent overall complication for the cohort was postoperative mental status change (13 patients [4.6%]), followed by hemorrhage (4 patients [1.4%]), seizure (4 patients [1.4%]), and hardware-related infection (3 patients [1.1%]). Mean LOS for all 284 patients was 1.19 ± 1.29 days (awake: 1.06 ± 0.46 days; asleep: 1.30 ± 1.67 days; p = 0.08). Overall, the 30-day readmission rate was 1.4% (1 awake patient, 3 asleep patients). There were no significant differences in complications, LOS, and 30-day readmissions between awake and asleep groups.

CONCLUSIONS Both awake and asleep DBS can be performed safely with low complication rates. The authors found no significant differences between the 2 procedure groups in adverse events, inpatient LOS, and 30-day readmission rates.

https://thejns.org/doi/abs/10.3171/2016.6.JNS152946

KEY WORDS adverse events; asleep deep brain stimulation; complications; deep brain stimulation; functional neurosurgery; intraoperative imaging; length of stay; readmission

Deep brain stimulation (DBS) is a well-established therapy for Parkinson’s disease and essential tremor, with its safety and efficacy having been demonstrated in multiple clinical trials. Several studies have shown relatively low rates of overall mortality (0%–8.3%), symptomatic intracranial hemorrhage (0.8%–5%), and infection (1.2%–15%) for DBS with microelectrode recording and intraoperative test stimulation. Traditionally, DBS has been performed with the patient “awake,” without general anesthesia, using intraoperative test stimulation with or without microelectrode recordings to guide lead placement. In contrast, “asleep” DBS is performed using either direct targeting (for the globus pallidus interna [GPI] and subthalamic nucleus [STN]) or indirect targeting (for the ventralis intermedius nucleus [VIM]), with the patient under general anesthesia and without intraoperative test stimulation. With improvements in MRI resolution and the accessibility of intraoperative imaging, asleep DBS surgery has gained popularity in the past several years.

As published reports on functional outcomes of asleep DBS increase, the question of whether there is a difference in how patients tolerate the procedure becomes pertinent. In particular, factors such as extended time under general anesthesia may influence perioperative outcomes in this population. However, no large studies have reported complication rates for asleep DBS, and relatively few series have analyzed length of stay (LOS) and readmission rates.
rates. As surgical techniques continuously evolve in an effort to improve the comfort, safety, and accessibility of the DBS operation, it becomes increasingly important to evaluate the impact of technical changes on perioperative safety, complication risk, and the duration of inpatient hospitalization. To address these issues, we retrospectively reviewed prospectively collected data on the adverse events, inpatient LOS, and 30-day readmissions for consecutive patients who had undergone either awake or asleep DBS surgery performed by a single neurosurgeon (F.A.P.).

Methods

Data were reviewed for all patients who had undergone either awake or asleep DBS surgery for movement disorders over a 3-year period (August 23, 2011–August 31, 2014). Targets were restricted to GPi, STN, and VIM (excluding one patient who underwent posterior subthalamic area DBS lead placement for essential tremor). Referring neurologists made all the primary diagnoses, and DBS candidacy was evaluated in a multidisciplinary consensus meeting of movement disorders neurologists, neurosurgeons, and neuropsychologists, at which targeting and laterality were also discussed. Both patient preference and the referring neurologist’s approval determined whether DBS was performed in the awake or asleep state. Informed consent was obtained that emphasized that asleep DBS does not involve intraoperative test stimulation and therefore constitutes off-label use since successful intraoperative test stimulation is a prerequisite in the US Food and Drug Administration labeling for DBS. All patients were followed up via a prospectively maintained database established for DBS patients of the senior author (F.A.P.). This database was approved by the Institutional Review Board at St. Joseph’s Hospital and Medical Center in Phoenix, Arizona. The most recent medical records of the patients were retrospectively reviewed to identify any additional complications not captured in the DBS database. All patients had a minimum of 6 months of follow-up.

Demographics and postoperative complications were reviewed. The total number of complications was analyzed based on the percentage of total leads. Hardware-related complications included infection, erosion, and revisions necessary for high impedance and lead malposition. Procedure-related complications included intraoperative and postoperative hemorrhage and seizure, cerebrospinal fluid leak, mental status change, pneumonia, and postoperative cerebral infarction. Inpatient LOS and 30-day readmission rates were also analyzed. For patients with multiple reasons for increased LOS, the primary etiology was noted.

Surgical Procedure

All procedures were performed utilizing uniform methods in each group (awake vs asleep). For all patients, preoperative imaging for direct (GPi or STN) or indirect (VIM) targeting was performed on a Siemens or General Electric 3-T MRI unit. DBS was performed with the patient either under intravenous sedation utilizing intraoperative test stimulation with or without microelectrode recordings (“awake”) or under general anesthesia without microelectrode recording or intraoperative test stimulation (“asleep”).

For awake patients, all leads were implanted using the Leksell (Elekta AB) stereotactic frame with microelectrode recordings to guide GPi and STN lead placement, and intraoperative test stimulation was performed to assess for clinical benefit and appropriate thresholds for side effects. An intraoperative CT (iCT) scan was obtained postoperatively on a CereTom or BodyTom mobile scanner (NeuroLogica Corp.) to document stereotactic accuracy, and image coregistration with the preoperative MRI was performed using FrameLink software (Medtronic, plc). Bilateral electrodes were typically placed in one operation.

Patients in the asleep group had the stereotactic Leksell frame or NexFrame (Medtronic, plc) bone fiducials placed in the operating room after the induction of general anesthesia. After DBS leads were implanted, an iCT scan was obtained before skin closure to assess accuracy and to determine the necessity for lead repositioning. If a lead was repositioned, an additional iCT scan was obtained to confirm accuracy.

Pulse generators were placed either on the same day as lead implantation or in a separate operation 1 week later. (In January 2014, our institution transitioned primarily to an “all-in-one” approach with the pulse generator and DBS leads implanted on the same day, rather than the traditional practice [in the US] of implanting the pulse generator 1–2 weeks after lead implantation.) We follow a strict protocol for pulse generator placement in which the targeted operative time is less than 30 minutes, only the primary surgeon handles the hardware, and exposure of the hardware is kept to a minimum (the scrub technician does not open or touch the hardware, and the generator is opened immediately before placing it in the pocket; this protocol was initiated following a personal communication with P. Starr on September 7, 2012, regarding protocols at the University of California, San Francisco). The generator is placed via a 5-cm incision 2 fingerbreadths beneath the clavicle. Both the retroauricular and the generator incisions are irrigated copiously with gentamicin before implantation, and vancomycin powder (500 mg) is placed into the retroauricular incision before closure.

In addition, beginning in August 2012, all patients underwent screening for methicillin-resistant Staphylococcus aureus (MRSA) before implantation. Patients and their household members were screened for MRSA at 4 sites: nares, throat, axilla, and groin. Patients who were MRSA positive were instructed to shampoo and shower with Hibiclens (chlorhexidine gluconate; Mölyneck Health Care) daily for 2 weeks before surgery. Seven days before surgery, patients initiated the application of mupirocin calcium 2% nasal ointment to both nares twice daily. Thirty minutes before skin incision, vancomycin (500 mg) was administered intravenously.

Statistical Analysis

Frequencies with percentages and means with standard deviations were used to describe the overall study cohort and patients with and without complications. Independent-
samples t-tests were used to compare mean differences for continuous variables, and chi-squared or Fisher’s exact tests were used for ordinal or dichotomous variables. To account for familywise error, we adjusted the significance threshold by dividing the standard cutoff of 0.05 by the number of statistical tests (50 tests) and obtained an adjusted threshold value of 0.001. Logistic regression models were then used to determine significant predictors of complications and mental status change, the most frequent complication. Because of the small sample size, patients with a dystonia diagnosis were excluded from regression analyses. We used SPSS Statistics for Windows, version 22 (IBM Corp.), for analyses.

**Results**

A total of 490 DBS electrodes were placed in 284 patients over a 36-month period. Demographics of the overall cohort and a comparison of the awake and asleep groups are shown in Table 1. Of the 284 patients undergoing implantation for DBS, 126 (44.4%) underwent awake surgery and 158 (55.6%) underwent asleep surgery. Pulse generators were placed either on the same day as lead implantation (29.6%) or in a separate operation 1 week later (70.4%). Mean follow-up was 21.4 ± 9.8 months (range 6–41 months). The most common indication for surgery was Parkinson’s disease (71.1%), followed by essential tremor (26.0%) and dystonia (2.8%).

**Awake Versus Asleep Group Demographics**

There were no differences in age or sex between the awake and asleep groups. There was a significantly greater proportion of bilateral cases in the asleep group (135 cases [85.4%]) than in the awake group (71 cases [56.3%]) (p < 0.001) and a greater proportion of staged cases in the awake group (102 cases [81.0%]) than in the asleep group (98 [62.0%]) (p = 0.001). Although the distribution of indications for surgery was significantly different between the asleep and awake groups (p < 0.001), the 2 groups consisted predominantly of Parkinson’s patients (awake 61.1%, asleep 79.1%), followed by essential tremor patients (awake 37.3%, asleep 17.1%) and dystonia patients (awake 1.6%, asleep 3.8%). The distribution of targets between the 2 groups was significantly different (p < 0.001). The VIM was the most frequent target in the awake group (52 cases [41.3%]), followed by the GPi (43 cases [34.1%]) and STN (31 cases [24.6%]). In the asleep group, the GPi was the most frequent target (94 cases [59.5%]), followed by the STN (36 cases [22.8%]) and VIM (28 cases [17.7%]).

There were no significant differences in the rest of the demographic variables between or within the awake and asleep groups. There were no intraoperative deaths, seizures, or aborted procedures. There was 1 perioperative death: a 69-year-old man with multiple medical comorbidities, who had undergone awake, staged, unilateral VIM electrode placement for Parkinson’s disease, died 2 months postoperatively as a result of gastrointestinal medical issues.

Subcategories of complications were compared between DBS techniques (awake vs asleep, staged vs all-in-one), number of leads placed (unilateral vs bilateral), and targets (GPi vs STN vs VIM) (Table 2). No patients experienced postoperative pneumonia, cerebrospinal fluid leak, or cerebral infarction (clinical or radiographic). Patients were returned to the operating room for infection, hardware failure, lead repositioning, and subdural hematoma due to a postoperative ground-level fall. Patients who experienced complications were significantly older than those who did not (68.2 ± 10.1 vs 63.6 ± 10.0 years, p = 0.03).

**Hardware-Related Complications**

The most common hardware-related complication was infection (3 patients [1.1%] with 6 leads [1.2%]). In all 3 cases, purulence developed at the pulse generator site only at the time of presentation. Wound cultures grew *Pseudomonas aeruginosa* and *Proteus mirabilis* (1 patient), coagulase-positive *S. aureus* and *Propionibacterium acnes* (1 patient), and MRSA (1 patient).

The second most frequent hardware complication was revision due to problems with impedance (2 patients [0.7%] with 3 leads). One case was detected immediately postoperatively in the recovery room, after an intraoperative normal impedance check. The patient was taken back to the operating room, where the problem was localized to the extension wire. It was switched out, and there were no further complications. The second patient also had normal impedance intraoperatively; however, at the initial programming 20 days after implantation, abnormal impedance was found on all combinations involving Contacts 3 and 1, ranging from 10,000 to 16,000 ohms. The patient underwent open interrogation 30 days after initial implan-

<table>
<thead>
<tr>
<th>Variable</th>
<th>All Patients</th>
<th>Awake Cohort</th>
<th>Asleep Cohort</th>
<th>p Value</th>
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<tbody>
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<td>No. of patients</td>
<td>284</td>
<td>126</td>
<td>158</td>
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<td>Mean age in yrs</td>
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<td>64.1 ± 10.5</td>
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</tr>
<tr>
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<td>181 (63.7)</td>
<td>85 (67.5)</td>
<td>96 (60.8)</td>
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<td>Female</td>
<td>103 (36.3)</td>
<td>41 (32.5)</td>
<td>62 (39.2)</td>
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<td>55 (43.7)</td>
<td>23 (14.6)</td>
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<td>Bilateral</td>
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<td>135 (85.4)</td>
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<td>PD</td>
<td>202 (71.1)</td>
<td>77 (61.1)</td>
<td>125 (79.1)</td>
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<td>Essential tremor</td>
<td>74 (26.1)</td>
<td>47 (37.3)</td>
<td>27 (17.1)</td>
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<td>Dystonia</td>
<td>8 (2.8)</td>
<td>2 (1.6)</td>
<td>6 (3.8)</td>
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<tr>
<td>Surgery type</td>
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<td>All-in-one</td>
<td>84 (29.6)</td>
<td>24 (19.0)</td>
<td>60 (38.0)</td>
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<tr>
<td>Staged</td>
<td>200 (70.4)</td>
<td>102 (81.0)</td>
<td>98 (62.0)</td>
<td>0.001</td>
</tr>
</tbody>
</table>

PD = Parkinson’s disease.
* Values represent number (percentage) or mean ± standard deviation.

**TABLE 1. Patient demographics and DBS categorization**

PD = Parkinson’s disease.

* Values represent number (percentage) or mean ± standard deviation.
Complications in awake versus asleep DBS

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tation, and the extension wire was replaced. There were no further complications.

Procedure-Related Complications

The most common procedure-related complication was mental status change (13 patients [4.6%] with 24 electrodes [4.9%]), followed by postoperative hemorrhage (4 patients [1.4%] with 7 electrodes [1.4%]). Of the 4 patients with hemorrhage, 1 suffered an asymptomatic acute subdural hematoma seen immediately on postoperative CT and 3 had normal postoperative CTs but developed symptomatic hemorrhages within 7 days of surgery. One patient experienced a focal subarachnoid hemorrhage, 1 had bilateral subdural hygromas, and 1 had both an acute subdural hematoma and an intraparenchymal contusion. The patient with unilateral subdural hematoma eventually required surgical evacuation on postoperative Day 14 because of symptomatic progressive mass effect (Fig. 1) and returned to neurological baseline postoperatively. Four patients (1.4%) with 6 electrodes (1.2%) had generalized tonic-clonic seizures; none had a seizure history. None of the 4 had new intracranial pathology when they again underwent CT scanning after the seizure.

Overall, the mean age of the patients with mental status change (70.2 ± 8.5 years) was not significantly higher than that of patients without such a change (63.7 ± 10.0 years) (p = 0.89). No statistically significant differences were found in categorical complication rates between awake and asleep, unilateral and bilateral, or staged and all-in-one groups or among the 3 DBS targets or 3 primary indications. There was, however, a somewhat higher percentage of patients in the all-in-one group who developed a change in mental status (p = 0.02). Multivariate logistic regression analysis demonstrated no association between complications and sex, age, primary diagnosis, and DBS technique (awake vs asleep, unilateral vs bilateral, or staged and all-in-one procedure). There was also no significant association among these variables with regard to mental status change, although patients who were 65 years old or older had an association trending toward significance (OR 4.71, 95% CI 0.99–22.32, p = 0.05) (Table 3). In the subset of 13 patients who experienced a mental status change, 9 had undergone asleep placement. In the asleep group, a significantly higher proportion of patients with mental status change were 65 years old or older (9 [100%] of 9) compared with those without mental status change (74 [49.7%] of 149) (p = 0.003).

Parkinson’s Disease Patients

Among the entire study cohort, 71.1% (202) of patients had Parkinson’s disease (38.1% awake, 61.9% asleep). A separate analysis was performed of hardware- and procedure-related complications in this group. Among the Parkinson’s patients, 10 patients (5.0%) with 16 leads (4.4% of 360 leads) experienced a mental status change, and more of these leads were placed using the all-in-one procedure (13 leads [11.2%]) versus the staged procedure (3 leads [1.2%]) (p = 0.009). No other differences were found among other technique variables (wake vs asleep, unilateral vs bilateral, GPi vs STN). Separate analyses for es-

| TABLE 2. Complications by diagnosis and DBS technique, leads, procedure, and target* |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Variable                        | Diagnosis       | p Value Unilat | Value Staged | Value All-in-One | p Value              | p Value              | p Value              |
| No. of electrodes               | PD              | 6 (1.2)         | 4 (0.7)       | 2 (0.5)         | >0.99               | 0.28               |
| Complication category           | Dystonia        | 1 (0.2)         | 1 (0.2)       | 0 (0.0)         | >0.99               | 0.41               |
| Infection                       | ET              | 3 (0.6)         | 2 (0.3)       | 2 (0.3)         | >0.99               | 0.37               |
| Erosion                         | Dystonia        | 3 (0.6)         | 2 (0.3)       | 2 (0.3)         | >0.99               | 0.37               |
| High impedance                  | PD              | 6 (1.2)         | 4 (0.7)       | 2 (0.5)         | >0.99               | 0.28               |
| Lead malposition                | Dystonia        | 7 (1.4)         | 5 (0.9)       | 2 (0.4)         | >0.99               | 0.28               |
| Procedure-related               | PD              | 7 (1.4)         | 7 (1.2)       | 3 (0.5)         | >0.99               | 0.28               |
| Hemorrhage                      | Dystonia        | 7 (1.4)         | 7 (1.2)       | 3 (0.5)         | >0.99               | 0.28               |
| Seizure                         | PD              | 7 (1.4)         | 7 (1.2)       | 3 (0.5)         | >0.99               | 0.28               |
| Mental status change            | Dystonia        | 7 (1.4)         | 7 (1.2)       | 3 (0.5)         | >0.99               | 0.28               |
| ET = essential tremor.          | PD              | 7 (1.4)         | 7 (1.2)       | 3 (0.5)         | >0.99               | 0.28               |
| * Values are number (percentage) unless indicated otherwise. The p values were determined by Fisher’s exact test. Statistical significance set at p < 0.01.
sentinal tremor and dystonia groups could not be performed because of their smaller sample sizes.

Length of Stay and Readmissions

The mean LOS for all 284 patients was 1.19 ± 1.29 days (range 1–19 days, median 1 day); for asleep patients it was 1.30 ± 1.67 days (range 1–19 days, median 1 day), and for awake patients it was 1.06 ± 0.46 days (range 1–6 days, median 1 day) (p = 0.08). At our institution, patients are typically admitted overnight and discharged the following morning, if appropriate. Sixteen patients (5.6%) required a prolonged hospital stay (≥ 2 nights) ranging from 2 to 19 days (mean 4.25 ± 4.21 days, median 3 days). The most common reasons for a prolonged inpatient stay were mental status change (4 cases), nausea/emesis (3 cases), and urinary retention (3 cases). Less common reasons were hemorrhage (2 cases) and oxygen desaturation, transient dysarthria, fever, and seizure (1 case each).

Four patients (1.4%) were readmitted within 30 days of implantation; all had undergone asleep DBS surgery. One patient had a mechanical fall and presented with bilateral subdural hematomas requiring evacuation. This patient required a prolonged hospitalization (13 nights) because of challenges with insurance authorization for acute rehabilitation. Of the remaining 3 patients, 1 developed spontaneous headache secondary to atraumatic bilateral subdural hygromas, which were conservatively managed with 2 nights of observation. The second patient was readmitted for concerns about seizure activity; however, seizure work-up was negative. This patient’s symptoms spontaneously resolved, and he was discharged after a 4-night hospitalization. The third patient was readmitted for new-onset seizure and was discharged after 1 night of hospitalization. The mean LOS for this group of patients upon readmission was 5.0 ± 5.8 days.

In the patient groups with prolonged hospital stays (≥ 2 or ≥ 3 nights) and 30-day readmissions, there were no statistically significant differences between asleep and awake groups, unilateral and bilateral leads, or staged and all-in-one procedures, nor were there differences among DBS targets or DBS indications. More patients in the asleep group (13 [8.2%]) than in the awake group (3 [2.4%]) (p = 0.04) had a hospital stay ≥ 2 nights, and more patients in the asleep group (8 [5.1%]) than in the awake group (1 [0.8%]) (p = 0.047) had a hospital stay ≥ 3 nights, although these findings did not reach statistical significance for this analysis (Table 4).

Discussion

There is growing interest in performing DBS under general anesthesia. Our initial experience has demonstrated equivalent functional outcomes between awake and asleep groups.\(^8\) We found that essential tremor patients who underwent asleep VIM placement demonstrate no difference in the percentage of postoperative improvement, as measured by their Bain and Findley Tremor Activities of Daily Living scores (48.6%),\(^8\) compared with awake patients (45.5%) (p = 0.35). Additionally, at 6-month follow-up, Parkinson’s patients who had undergone asleep GPi placement had significant improvement in mean off-medication motor scores (48.4 vs 29.8, p < 0.001) and in 39-item Parkinson’s Disease Questionnaire (PDQ-39) scores (50.3 vs 42.0, p = 0.03). Daily levodopa-equivalent doses were also significantly decreased at 6 months (1207 vs 1035 mg, p = 0.004).\(^27\) As reports of functional outcome data for asleep DBS have increased in the medical literature, the effect of anesthesia on perioperative recovery has become an area of growing concern for neurologists and patients. Complication rates and LOS data for a large series of patients who have undergone asleep DBS surgery under general anesthesia have not been reported. In this study, we found no difference in hardware- or procedure-related complications in patients undergoing asleep and awake surgeries. In addition, we found no difference in the mean LOS of these 2 groups.

Complications

Our overall hardware complication rate was 6.0%, which is within the lower range reported for previous studies (4.9%–22.8%).\(^3,6,7,9,15,16,19,20,29,32,33,35,42,43\) Our most frequent hardware-related complication was infection (6 leads [1.2%] in 3 patients [1.1%]), and this rate is also within the lower range of previously published infection rates (0.4%–15.2%).\(^5,7,10,11,12,15,16,19,20,29,32,33,35,42,43\) This wide range

<table>
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<th>Variable</th>
<th>Complications</th>
<th>Mental Status Change</th>
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<tr>
<td>Sex</td>
<td>p</td>
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<tr>
<td>Asleep vs awake</td>
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<tr>
<td>Unilateral vs bilateral</td>
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<td>PD vs ET</td>
<td>0.52</td>
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<tr>
<td>Age ≥65 yrs vs &lt;65 yrs</td>
<td>0.33</td>
<td>0.56</td>
</tr>
<tr>
<td>All-in-one vs staged</td>
<td>0.16</td>
<td>1.90</td>
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</table>
of reported infection rates includes some smaller series of 30–40 patients.\textsuperscript{19,22,37} However, 2 recent large, single-center series have also demonstrated low infection rates of 4.5% in 759 leads placed in 420 patients\textsuperscript{38} and of 1.18% in 510 leads placed in 392 patients.\textsuperscript{32}

We believe that our low infection rate is primarily attributable to strict adherence to our pulse generator placement protocol, which includes restricting generator handling to one person, applying vancomycin powder to the cranial incision, and using a preoperative MRSA-screening and treatment protocol. We used the same generator protocol across all patients and found no difference in hardware-related complications in asleep versus awake groups, unilateral versus bilateral groups, or all-in-one versus staged groups, or among DBS target groups.

Mental status change was the most common procedure-related complication (24 leads [4.9%] in 13 patients [4.6%]), which is consistent with data in the literature, with reported rates of 4.7%–27.7% for unilateral procedures, 10.7%–14.6% for simultaneous bilateral procedures, and 3.5%–22% for staged bilateral procedures.\textsuperscript{32} The rate of mental status change in our patients with simultaneous bilateral implanted electrodes (5.4%) is lower than historical rates (10.7%–14.6%).\textsuperscript{16,19,32,43} This finding may be related, in part, to the incomplete capture of all patients experiencing a mental status change, because of the variability and subjectivity of symptoms within this category and the challenges faced when discerning symptoms from a chart review. It may also be related to the relative decrease in time required to place a second electrode (data not reported).

Hemorrhage occurred in 4 patients (1.4%) with 7 leads (1.4%). Three patients were symptomatic, and 1 of these 3 required additional craniotomy for subdural hematoma evacuation. This rate is within the range of previously reported rates for intracranial hemorraghes (1.1%–6.29%) in studies with 30–420 patients.\textsuperscript{3,6,7,9,15,16,19,20,29,32,33,35,42,43} Multivariate analysis showed no increased risk in overall complications associated with sex, increased age (\( \geq 65 \) vs < 65 years), diagnosis (Parkinson’s disease vs essential tremor), or DBS technique (awake vs asleep, unilateral vs bilateral, all-in-one vs staged). However, further analysis of the asleep group showed that a significantly higher proportion of patients with than without mental status change were 65 years old or older. This finding is in line with established knowledge that older patients are more likely to experience mental status changes after general anesthesia, regardless of the surgical procedure. All mental status changes in our patients were temporary, and patients were discharged home at their neurological baseline.

Previous studies have also demonstrated a significant association between increasing age and hemorrhage, raising concern that older patients may have a higher complication rate and may not tolerate DBS surgery as well as younger patients.\textsuperscript{45} However, 2 recent database analyses of 661 patients undergoing DBS for essential tremor and of 1757 patients undergoing DBS for Parkinson’s disease with 90-day follow-up also demonstrated no increased risk of complications associated with increasing age from < 50 to 90 years for either group.\textsuperscript{40,41} Further subanalysis of patients experiencing mental status change showed a trend toward significance for a positive association with those 65

<table>
<thead>
<tr>
<th>TABLE 4. Length of stay and 30-day readmissions by number of patients and DBS technique, leads, procedure, and target*</th>
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</thead>
<tbody>
<tr>
<td><strong>Variable</strong></td>
</tr>
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<td>No. of patients</td>
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<td>Hospital stay ≥2 nights</td>
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<tr>
<td>Hospital stay ≥3 nights</td>
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<td>Readmit w/in 30 days</td>
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* Values are number (percentage) unless indicated otherwise. The p values were determined by chi-squared test. Statistical significance set at p < 0.01.
years old or older; however, a larger sample size is likely needed to detect true significance.

We performed a separate analysis among the Parkinson’s patients (202 patients) to eliminate variability introduced by other disease groups (essential tremor and dystonia). Results showed that significantly more patients experienced mental status change in the all-in-one placement group versus the staged placement group (p = 0.009). This finding may be related to the longer duration of continuous anesthesia for patients undergoing the all-in-one procedure.

Length of Stay and Readmission Rates

There are limited data on LOS and readmission rates after DBS. While most patients stay only 1 hospital night after DBS implantation, a prolonged LOS can occur for several reasons. In a recent series by Mikos et al.,24 115 unilateral DBS electrodes were placed in Parkinson’s patients, 21% of whom required a hospital stay of more than 1 night. The most common reason for an increased LOS was mental status change, followed by hemorrhage, with a total LOS ranging from 2 to 60 days. In Goodman et al.’s30 series of 191 electrodes in 100 patients, the mean LOS was 3.1 days; however, these patients were admitted 1 day before surgery to facilitate preoperative transitioning off Parkinson medications. Mental status change was also the most frequent cause of increased LOS in this group. In our series, only 5.6% of patients required a hospital stay of ≥ 2 nights, and the mean LOS was 1.19 ± 1.29 days. Our patients are admitted on the day of surgery, rather than on the preceding day.

The leading cause of increased LOS in our patients was also mental status change (4 patients), followed by nausea and/or emesis (3 patients) and urinary retention (3 patients). All 3 patients with an increased LOS due to nausea and/or emesis and 2 of the 3 patients with urinary retention were in the asleep group. Although there was a trend toward more asleep patients with a ≥ 2-night hospital stay (8.2%) than awake patients (2.4%) (p = 0.04) and more asleep patients with a ≥ 3-night hospital stay (5.1%) than awake patients (0.8%) (p = 0.047), these findings did not reach statistical significance for this analysis. Given the standard threshold of p < 0.05, the comparison between 13 patients (8.2%) and 3 patients (2.4%) would be considered statistically significant, but with only 53% power. To account for familywise error, we adjusted the threshold for our study to p < 0.01, and thus the association between hospital LOS ≥ 2 and ≥ 3 days with awake versus asleep implantation was not statistically significant. Setting alpha at 0.01 would require at least 200 patients in the awake group and 251 in the asleep group to detect a statistically significant difference between 8.2% and 2.4%. All patients with a ≥ 3-night hospital stay (9 patients [3.2%]) had Parkinson’s disease, although this finding was not statistically significant either. These trends may be related to factors associated with the postoperative side effects of general anesthesia. Multivariate analysis could not be performed because of the small sample size.

A recent study reviewing 30-day readmissions for cranial neurosurgical procedures within the categories of neoplasm, vascular disease, seizure, and trauma found readmission rates that ranged from 14% (seizure group) to 24% (vascular group).28 In a series of 303 patients from our institution, the 30-day readmission rate after transphenoidal surgery for pituitary tumor was 8.9%, with delayed hyponatremia (55.6%) as the most frequent cause of readmission.4 Relatively few studies have evaluated readmission rates for DBS patients; however, a recent study by Jacob et al.18 found a 30-day readmission rate of 4.3% for 211 DBS patients. In the current study, we found a comparably low 30-day readmission rate of 1.4%. The lower readmission rates for DBS patients compared with those for other neurosurgical patients probably reflect the nature of the pathology being treated and the increased severity of illness among patients with neoplastic, vascular, epileptic, and traumatic diseases. These neurosurgical pathologies often necessitate emergent or urgent surgical intervention, whereas DBS procedures are performed on an elective basis.

Avoidance of Complications and Lessons Learned

Infection

The 3 patients with hardware-related infections required complete explantation despite attempts to salvage the DBS system. To avoid futile salvage attempts in the future, we therefore now explain to patients preoperatively that the definitive treatment for infection is complete system explantation. Interestingly, these 3 infections occurred early in both the awake experience (Case 1) and the asleep experience (Cases 4 and 7). This suggests that, with appropriate attention and experience, the rate of infection for DBS surgery can be kept to a minimum. After a MRSA infection occurred in a 52-year-old male patient, we instituted a MRSA-screening program. We also use vancomycin powder (1000 mg) during lead placement for the frontal incisions, vancomycin powder (500 mg) during battery placement for the retroauricular incision, and a TYRX (Medtronic, plc) antibacterial pouch for the generator, as well as the aforementioned protocol regarding surgery duration and hardware handling.

We believe that the MRSA-screening process is effective, because there were no additional MRSA infections after its initiation (August 2012) through the end of the study period (August 2014). However, other factors outlined in the pulse generator protocol, such as the use of vancomycin powder, may have also contributed to this finding. Overall, 225 patients were swabbed during the study period, and 6 patients (2.67%) tested positive preoperatively and were treated. Four patients were colonized in the nares; 1 patient in the throat; and 1 patient in the nares, axilla, and throat.

On the basis of recommendations from our Infectious Disease Department after reviewing the initial MRSA detection rate, we modified our protocol (as of December 2015) to swab patients who answer “yes” to one of our screening questions: Have you or any of your family members been treated for MRSA? Have you or any of your family members worked in a health care setting? Have you had multiple hospitalizations? Have you been a carrier of MRSA? We also ask patients 2 weeks before their scheduled surgery about any antibiotic use in the last 3 months. In light of the potentially severe consequences of hardware infection and the fact
that MRSA screening is relatively inexpensive and easy to perform, we believe that it is worthwhile to pursue such screening.

Intraoperative Impedance Checks

Impedance checks can be used to verify the appropriate overlap of contacts and to identify fractures in the lead extension system. We now check impedance 1) after securing all connections with the torque wrench to verify appropriate connectivity; 2) after placing and tucking all hardware pieces in their respective pockets to verify that the hardware was not injured by manipulation; and 3) after final closure to verify no nicking by sutures. We have found abnormal impedance in a number of extension wires at the second check after finding normal impedance at the first check and were able to replace these extensions before closure.

Seizure

The mechanism of seizure can be venous congestion at the cortex, which can be a consequence of electrocautery. Like others, we believe that it is necessary to open the pia before introducing the cannula (personal communication, K. Foote, June 6, 2012). We have been able to successfully minimize pneumocephalus by opening the dura with monopolar electrocautery and a sharp tip stylet. We then insert the stylet into the pia to create an opening without the use of electrocautery. Fibrin glue is placed around the cannula to prevent cerebrospinal fluid loss and subsequent brain shift before lead placement. Since adopting this technique, we seldom directly visualize the brain. Instead, key steps are the use of preoperative MRI with contrast, trajectory selection that avoids sulci or blood vessels, and strict adherence to the trajectory coordinates on the frame (that is, the ring and arc coordinates on the Leksell frame) to allow for “blind” penetration of the brain in a safe manner.

To account for gravitational shift, we also maintain the same patient positioning (that is, the patient supine with the head of the bed flat) during MRI and CT scanning and throughout the procedure for asleep surgeries. Our asleep placement protocol transitioned from frameless to frame-based placement in January 2013, when we began keeping the head of the bed at 0° for MRI and CT scanning and for the duration of the procedure. After this transition, significant improvement in accuracy was demonstrated in a recent multivariate analysis. Increasingly, for our awake cases, the head of the bed is also kept flat for microelectrode recording and lead placement, and the head angle is adjusted upward only for test stimulation.

Study Limitations

There are several limitations to this study, including its retrospective nature and relatively short duration of follow-up. The mean follow-up is 31.9 months. Our sample size was 214. ± 9.8 months. Longer-term follow-up would have allowed us to capture more adverse events, potentially increasing the likelihood of detecting a significant difference among groups. Nevertheless, because of our small sample size and the low complication rates in each adverse event category, the detection of statistically significant differences among groups is unlikely. The ability to perform additional multivariate analysis beyond what was included is also limited. Twenty-four leads were associated with the most common complication (mental status change). As shown in Table 2, the p value associated with this comparison is 0.29, equating to an achieved power of 24%. Given the same proportions, a sample size of 3117—more than 6 times the size of our sample—would be required to achieve significance. Similarly, the p value associated with seizures was 0.23, with an achieved power of 20%. Given these same proportions, 2490 patients would be required to achieve 80% power.

Another possible limitation is our restricted ability to capture accurate readmission rates. We were able to access only the 30-day readmission data for patients who were readmitted either to our institution or to other hospitals that sent their medical records to our offices. There may have been patients readmitted to other hospitals about whom we were never notified, such as patients referred from out of town, although we suspect that this number would be quite small since most of our patients reside locally.

A further limitation is the slight variability in patient characteristics between and within the awake and asleep groups. There were a significantly higher proportion of essential tremor cases in the awake versus the asleep group and a significantly greater number of GPi leads in the asleep versus the awake group. The asleep group had a greater number of bilateral cases and all-in-one cases. The remaining variables were not significantly different between and within the awake and asleep groups. We recognize that these few variations are potential confounders to the analysis; however, we do not believe that they significantly affect our ability to compare the awake and asleep groups. Because of the overall low rate of complications, we analyzed the cohort as a whole to capture all the potential adverse events.

The generalizability of our results may be limited since our specific institutional protocols for awake versus asleep DBS surgery, generator placement, and preoperative MRSA screening may not be standard at other institutions. Furthermore, given the overall low rate of complications captured and the power limitations with regard to sample size, these results may not be readily generalizable. Nevertheless, as asleep DBS surgery becomes more common, this study will provide important initial data on the rates of adverse events, LOS, and postoperative readmission rates in a relatively large series of patients undergoing DBS surgery with general anesthesia compared with data from patients undergoing traditional awake DBS surgery with microelectrode recording.

Conclusions

Both awake and asleep DBS surgery can be performed safely with a low rate of complications. LOS and 30-day readmission rates are low for both groups. In our initial experience, asleep surgery versus traditional awake surgery probably confers no difference in the incidence of complications, LOS, or 30-day readmissions. Patients in the asleep group who were 65 years old or older may have had an increased likelihood of postoperative mental status change due to the effects of general anesthesia.
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Disclosures
Dr. Ponce is a consultant for Medtronic, plc, and has received financial support from the Barrow Center for Neuromodulation.

Author Contributions
Conception and design: Ponce, Chen, Mirzadeh. Acquisition of data: Ponce, Chen, Lambert. Analysis and interpretation of data: Chen. Drafting the article: Ponce, Chen. Critically revising the article: Ponce. Reviewed submitted version of manuscript: Ponce. Statistical analysis: Chen, Chapple. Administrative/technical/material support: Lambert. Study supervision: Ponce.

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