Intrathecal Baclofen for Severe Spasticity: Longitudinal Data From the Product Surveillance Registry

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Objective: To assist in the assessment of intrathecal baclofen (ITB) therapy risks and benefits by providing surgical intervention rate, safety, and elective device replacement rate data.

Materials and Methods: An ongoing prospective, long-term, multicenter Product Surveillance Registry (PSR) (NCT01524276) enrolled consented patients implanted with the SynchroMed II infusion system. Pump and catheter performance data were collected, with patients followed prospectively for events related to the device, procedure, and therapy. Investigators provided event descriptions, patient symptoms, and patient outcomes.

Results: We analyzed registry data from 1743 patients (77% adult, 46.8% female) treated with ITB for severe spasticity at 53 registry sites between August 2003 and October 2017, for an accumulated 6481 patient-years. Discontinuation from the registry was largely (58.6% of discontinued patients) due to study site closure and patient relocation; exit due to an adverse event was limited to 0.3%. After 10 years, 87.2% of adult and 76.3% of pediatric patients continued with ITB. Overall, 99.1% of pumps reaching end of battery life were replaced at the time of explant.

Conclusions: ITB therapy for the treatment of severe spasticity requires surgical implantation of a programmable infusion system for chronic drug delivery. If complications arise, many necessitate surgical intervention for correction. For spinal and cerebral spasticity in pediatric and adult patients, discontinuation rates due to an adverse event were low (0.3%), and there was high acceptance (99.1%) of surgical intervention for therapy continuation. Patient/caregiver willingness to accept surgical and other risks for therapy continuation was extremely high.

Keywords: intrathecal baclofen, registry, risk, spasticity

Conflict of Interest: The Product Surveillance Registry is maintained by Medtronic, Inc. Authors Spencer, Stromberg, and Weaver are employees of Medtronic. The other authors received financial support for this research from Medtronic. The author(s) report receiving financial support for the research. Funding for the Product Surveillance Registry, data monitoring, statistical analysis, and medical writing were provided by Medtronic, Inc.

INTRODUCTION

Severe spastic tone can be the most disabling consequence of a neurologic insult that creates an upper motor neuron lesion from either a cerebral (e.g., cerebral palsy, acute brain injury, stroke) or spinal (e.g., spinal cord injury, multiple sclerosis) etiology. Spasticity is clinically recognized as a disorder of sensory-motor control brought about by an imbalance of supraspinal descending excitatory/inhibitory neurons that regulate muscle tone (1).

Treatment has focused on improving function, mobility, and range of motion to alleviate pain and prevent secondary complications. Conventional medical management (CMM) involves physical therapy and antispasmodic medications, with baclofen the most frequently used. The effectiveness of oral baclofen is often limited by its systemic side effects of dizziness, sedation, confusion, lethargy, and muscle weakness (2). Intrathecal baclofen (ITB) administration delivers drug directly to receptor sites, reducing
ITB Registry Longitudinal Data

MATERIALS AND METHODS

We analyzed data from the Product Surveillance Registry (PSR) to assess the safety (adverse events or AEs related to the device, implantation, or infusion therapy), durability (duration of pump implants and patient survival from study exit), and patient satisfaction (elective pump replacement) with ITB therapy for spasticity. The PSR comprises the largest, multicenter cohort of ITB-treated patients worldwide, covering an accumulated 6481 patient-years of ITB experience.

Registry Description

The Medtronic Implantable Systems Performance Registry (ISPR; ClinicalTrials.gov Identifier: NCT01524276), initiated in 2003, is described in detail by Konrad et al. (8). Results presented there include data collected under the ISPR protocol, which focused on a single manufacturer’s product performance at a number of implant centers around the world in which longitudinal patient follow-up was highly likely for the life of the implant. In 2013, the registry expanded data collection with a corresponding name change to the PSR. The PSR platform was designed to conduct ongoing nonrandomized, active prospective post-market surveillance under a common protocol, with specific appendices for neuromodulation products/therapies, by enrolling patients with an eligible product—in this case, implanted ITB pumps. Prior to patient enrollment, all sites obtained Ethics Committee/Institutional Review Board approval to allow tracking of the device performance in each consented patient. The PSR sites contributing to these data are noted in the Acknowledgments section.

Patients

Potential PSR patients were identified from the practices of participating physicians as meeting the specific indication of severe spasticity for the SynchroMed II infusion system (Medtronic, Inc. Minneapolis, MN, USA) and were enrolled at initial implant or at the time of pump replacement; excluded were patients enrolled/implanted for the treatment of chronic nonmalignant pain and cancer pain. The patient or legally authorized representative provided written authorization and/or consent per institution and geographical requirements prior to data collection. Patients inaccessible for follow-up, excluded per local law, or currently enrolled in or planning to enroll in any concurrent drug and/or device study that may confound results were excluded. Data were only included for patients who consent to enroll. After enrollment, patients were followed longitudinally per standard of care, with status updates obtained every 6 months with no predefined duration. Data for all global registry centers and all patients enrolled are reported.

Data Collection

The registry was created to monitor the performance of the implanted infusion systems, but the types of data collected have evolved over time. Safety data were originally focused on device-related events but expanded in 2010 to include all serious AEs and all events related to implanted or external components of the infusion system, the implant procedure, or the infusion therapy. Reporting of device-related surgical intervention, however, has remained consistent throughout the course of the registry. Collection of outcome data by registry sites (i.e., EQ-5D, numerical pain rating scale) was added to the PSR in 2013; however, a spasticity or tone assessment was not added. Elective pump replacement was assessed here as a surrogate for patient satisfaction. All data were entered by site personnel in a centralized database.

Analytic Methods

Data included in this analysis were collected from August 2003 through October 31, 2017 from patients in the ongoing PSR who were treated with ITB for severe spasticity. Summary statistics are presented either as percentages for categorical variables or as mean (standard deviation, SD; or minimum/maximum) for continuous values.

In order to evaluate the ongoing risk faced by ITB patients, the rate of surgical interventions by year post-implant was calculated as follows: number of nonbattery replacement surgical interventions per number of active patients within each year post-implant for the subset of patients who were enrolled in the registry with their initial infusion system.

Long-term therapy retention was assessed using time-to-event Kaplan-Meier survival estimates. Patients with their initial implant at the time of enrollment were included in the analysis. Study exit due to an AE or device event or due to an inactive system (i.e., therapy abandoned for greater than 6 months or system explanted without replacement) was the event of interest. Survival time (i.e., duration of therapy retention) was defined as months from implant to therapy discontinuation. Patients who were active in the registry or who exited from the registry due to nontherapy discontinuation reasons were censored at their last visit.

RESULTS

A total of 7975 pump patients were enrolled at 64 sites across the US, Europe, and Latin America, with 1743 (46.8% female) from 53 sites being treated for severe spasticity. The majority of patients with spasticity (77%, 1350/1743) were at least 18 years old.
old at enrollment. Most of the pediatric patients (patients under 18 years old at enrollment) were implanted for the treatment of cerebral palsy (CP, 78%, 307/393). Among adults, the most common treatment indication was multiple sclerosis (MS, 33%, 446/1350), followed by spinal cord injury (SCI, 17%, 231/1350). Figure 1 provides spasticity etiology for adult and pediatric patients. At the time of enrollment, 72% (1255/1743) of patients were enrolled with their initial pump and were naïve to infusion therapy. The remaining 28% (488/1743) of patients were enrolled with a pump replacement. For patients with an initial implant, the average age for the adult cohort was 46 years (±14, 18-86) and for children 10 years (±4, 2-18).

Patient participation within this registry is not defined by any endpoint date or duration of follow-up. Patients remained in the registry until they were no longer available for follow-up (e.g., study site closure, lost to follow-up, change to another treating institution or health-care provider) or their continued participation is no longer warranted (e.g., system explant or therapy discontinuation). Over time there have been 1202 spasticity patients (69%) who discontinued from the registry, with the reasons for study exit for these patients in Table 1. “Registry study site closure” (29.7%) and “Patient care transferred” (28.9%) account for 704 (58.6%) of these discontinuations, with “Lost to follow-up” and “Patient no longer available for follow-up” accounting for another 112 (9.3%) patients. Therapy events (e.g., surgical interventions) for these 816 enrolled patients were included through the date of their registry discontinuation, but final outcome remains unknown. Overall, the average duration of patient follow-up was 44.6 months (SD = 40.2) and ranged from 0 months to more than 13 years, with an accumulated follow-up of 6481 patient-years. Unlike studies with a predefined finite duration, this registry has been active for more than 16 years and continues to follow existing patients and enroll new patients.

Table 2 summarizes the number of surgical device interventions (i.e., pump or catheter explant or surgical procedure to modify an implanted component, such as pump repositioning or catheter revision) by year postimplantation for patients who enrolled with an initial implant in the registry (therapy naïve). A count of the number of interventions, the number of patients (therapy naïve) who were active during each year, and the rate of intervention/patient was summarized by age group and overall. The table excludes pump explants due to normal battery depletion, which had no associated AE. Each period reported included interventions occurring prior to the last year reported, that is, Year 0-1 included surgical interventions that occurred before completing the first year of implant, Year 1-2 included surgical interventions that occurred on or after 1 year of implant, but prior to the second year of implant, and so on.

Study exit reasons were used as a surrogate indicator of patient satisfaction with ITB therapy. A patient who discontinued from the registry due to an adverse or device event or due to an inactive system was considered a discontinuation from the registry due to dissatisfaction with the therapy. Patient survival from therapy-related discontinuation is summarized for adult and pediatric patients in Fig. 2. Within the cohort of patients receiving an initial pump implant at enrollment, 87.2% (95% confidence interval [CI]: 83.4%, 90.1%) of adult patients and 76.3% (95% CI: 68.6%, 82.4%) of pediatric patients maintained ITB therapy (i.e., had not discontinued from the registry for reasons related to adverse or device events or therapy discontinuation) through at least 10 years. Patient survival from therapy-related discontinuation is shown by spasticity indication in Fig. 2.

Implanted, battery powered pumps eventually require replacement due to battery depletion. In the case of the SynchroMed II infusion system, the designed performance life is capped at 84 months, with a typical time to replacement of 75 months for the currently available infusion system. The clinician-programmer displays an anticipated end of service (EOS) date with each pump interrogation; and an elective replacement indicator (ERI) message is displayed 90 days in advance of the EOS. This lead time for a necessary surgical intervention to continue ITB therapy, in contrast to the more acute decisions needed in cases of system complications, allows for a thorough patient/caregiver assessment of therapy benefits in deciding whether to undergo the replacement procedure. A total of 426 patients progressed to the point of at least one elective pump replacement (462 total pumps: 364 in ITB naïve patients, 98 in patients enrolled with a pump replacement) within the registry; 32 patients had undergone more than one pump replaced while enrolled in the registry. Mean implant duration (months) is summarized in Table 3 for the 462 pumps explanted due to a normal battery depletion. More than 99% of the 462 pumps that were explanted due to normal battery depletion were replaced on the same day, with four patients (4/364 ITB naïve at enrollment) electing system explant; two due to loss of therapeutic benefit and one each due to “patient request” and

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**Figure 1.** Spasticity etiology by adult/pediatric patients. [Color figure can be viewed at wileyonlinelibrary.com]
Table 1. Study Exit Reasons.

| Exit reason                              | Adult patients | Pediatric patients | Total** |
|------------------------------------------|----------------|-------------------|---------|
| Adverse event or device event            | 2 (0.2%)       | 2 (0.7%)          | 4 (0.3%)|
| Therapy discontinued                      | 70 (7.6%)      | 54 (19.6%)        | 124 (10.3%)|
| Death                                    | 131 (14.1%)    | 47 (17.0%)        | 178 (14.8%)*|
| Eligibility criteria not met†            | 11 (1.2%)      | 0 (0.0%)          | 11 (0.9%)|
| Patient withdrawal of consent            | 19 (2.1%)      | 7 (2.5%)          | 26 (2.2%)|
| Withdrawal of patient by physician‡      | 16 (1.7%)      | 20 (7.2%)         | 36 (3.0%)|
| Other                                    | 6 (0.6%)       | 1 (0.4%)          | 7 (0.6%)|
| Final status unknown: Follow-up terminated | 54 (5.8%)     | 11 (4.0%)         | 65 (5.4%)|
| Lost to follow-up                        | 282 (30.5%)    | 65 (23.6%)        | 347 (28.9%)|
| Patient no longer available for follow-up| 43 (4.6%)      | 4 (1.4%)          | 47 (3.9%)|
| Site closure§                            | 292 (31.5%)    | 65 (23.6%)        | 357 (29.7%)|
| Total                                    | 926            | 276               | 1202    |

*Reason for study exit was reported as death for 178 patients, 169 of which were reported as unrelated to the device. Patient indications for device implants in those who died were multiple sclerosis (51), cerebral palsy (41), spinal cord injury (22), stroke (10), traumatic brain injury (6); the remainder (39) were in patients with other/unknown primary intrathecal baclofen indication. Two deaths involved pump pocket infections in immunocompromised patients, which are not classified as device-related. One patient treated with ITB for traumatic brain injury-related spasticity experienced a pump pocket infection within one month of implantation that progressed to sepsis and resulted in death; the second patient, being treated for CP-related spasticity, experienced pump pocket erosion and an MSSA-positive infection two years postimplantation that resulted in death. Relationship to device/therapy was “unknown” in an additional seven patients: four multiple sclerosis and three cerebral palsy.
†Includes situations such as consented/enrolled before implant but pump implant not completed.
‡Transfer of care for enrolled patients is typical for pediatric patients transitioning to adult care and in cases of patient relocation.
§Over the 15-year span for ISPR/PSR a total of 53 sites enrolled patients implanted for the treatment of spasticity with 27 active sites through October 2017. Site closure takes place primarily due to principal investigator retirement or relocation.
**Twenty-five percent of these patients were enrolled with a replacement pump.

“unacceptable follow-up and maintenance requirements” (Table 3).

DISCUSSION

Our safety analysis included 1743 patients with average duration of patient follow-up of 44.6 months (SD = 40.2) that ranged from 0 months to more than 13 years, and an accumulated 6481 patient-years of experience. The number of surgical interventions per year post-implant for therapy naïve patients was highest in the first year (113 total) followed by year 6-7 (63 total) and year 3-4 (42 total). Dura-

Table 2. Surgical Interventions by Year Postimplant, by Age Group (Initial Implant).

| Year postimplant | No. of interventions | No. of patients | Intervention/patient | No. of interventions | No. of patients | Intervention/patient | No. of interventions | No. of patients | Intervention/patient |
|------------------|----------------------|-----------------|----------------------|----------------------|-----------------|----------------------|----------------------|-----------------|----------------------|
| 0-1              | 83                   | 738             | 0.112                | 30                   | 224             | 0.134                | 113                   | 962             | 0.117                |
| 1-2              | 25                   | 592             | 0.042                | 18                   | 215             | 0.084                | 43                    | 807             | 0.053                |
| 2-3              | 25                   | 537             | 0.047                | 14                   | 216             | 0.065                | 39                    | 753             | 0.052                |
| 3-4              | 28                   | 470             | 0.060                | 14                   | 192             | 0.073                | 42                    | 662             | 0.063                |
| 4-5              | 22                   | 391             | 0.056                | 17                   | 177             | 0.096                | 39                    | 568             | 0.069                |
| 5-6              | 22                   | 332             | 0.066                | 13                   | 149             | 0.087                | 35                    | 481             | 0.073                |
| 6-7              | 30                   | 275             | 0.109                | 33                   | 125             | 0.264                | 63                    | 400             | 0.158                |
| 7-8              | 12                   | 207             | 0.058                | 1                    | 103             | 0.010                | 13                    | 310             | 0.042                |
| 8-9              | 5                    | 152             | 0.033                | 8                    | 83              | 0.096                | 13                    | 235             | 0.055                |
| 9-10             | 5                    | 107             | 0.047                | 3                    | 70              | 0.043                | 8                    | 177             | 0.045                |

ITB therapy was originally evaluated for the treatment of severe spasticity of spinal origin (9,10), with indications expanded to include severe spasticity of cerebral origin, including cerebral palsy (11), brain injury (12), and stroke (13-15). Previous studies have demonstrated efficacy in spasticity of multiple origins, with long-term benefits (11,16-18). A 1997 meta-analysis of 27 studies with 490 ITB-treated patients included spasticity due to SCI (42% of patients), MS (33%), CP (12%), and other causes (13%) (19). The results demonstrated significant decreases in mean Ashworth and spasm scores in each diagnostic group at 18 months average follow-up; 8% of implanted patients discontinued treatment by 1 year in these early studies.

More recently, Cramer et al. compared the effectiveness of ITB therapy with CMM in 60 subjects with post-stroke spasticity in a randomized multicenter study (15). In addition to improvements in...
spastic hypertonia and muscle tone, a positive trend toward greater functional ability favored ITB. AEs (collected regardless of therapy relatedness) were more frequently reported in the ITB group, where 24/25 (96%) of subjects reported 149 events, compared with CMM, where 22/35 (63%) subjects reported 77 events. In the ITB group, approximately half of all events (69 of 149) occurred during the initial six-week dose-titration period. Reported events were consistent with the known safety profile of ITB. Four events in four patients randomized to the ITB arm were classified as serious adverse device events (device dislocation, device occlusion, implant-site infection and intracranial hypotension), and three led to device revision.

These randomized controlled trials and prospective case series include a limited number of participants enrolled and managed within tight study protocols for limited duration, and thus run the risk of under- or overestimating some therapy-related AEs. Real-world evidence from a large registry provides additional data to offer patients and clinicians a clearer assessment of the safety profile of a therapy. The PSR is the largest implanted drug delivery system registry in the world, with global and diverse clinical-site participation that offers a unique and pragmatic perspective on long-term ITB and adds to the general applicability of reported outcomes. Although spasticity-specific clinical outcomes in the PSR may be limited, therapy retention and elective pump replacement may reflect overall patient satisfaction and continued commitment to ITB therapy. The registry also selects study sites that represent commitment for long-term follow-up of patients and therefore will likely reflect more experienced and optimal care and management of the device when compared with novice implant centers or centers that implant ITB occasionally. Therefore, our AE rate may underestimate the complication rate and overestimate satisfaction compared with implant sites and subjects reported elsewhere in the literature.

AEs within the PSR are collected prospectively with risk-based monitoring to ensure accuracy and completeness. The risks associated with ITB therapy and the SynchroMed infusion system have been well studied, with publications on individual patient cases or

Figure 2. Patient survival from therapy-related study exit: a. By indication; b. by age category. [Color figure can be viewed at wileyonlinelibrary.com]
series reports (20–23) and as many as 359 patients (24). The 1743 patients enrolled in the PSR and their average 44.6 months and cumulative 6481 patient-years of follow-up provide a currently unmatched source of safety data. Although the individual rates of specific complications are important in assessing safety, presenting “surgical risk” with a therapy for which surgical revision is often required for AE resolution has more relevance to both the treating physician and patient. As presented here, the risk of surgical revision (number of interventions/ITB naïve patients followed) is highest (adult: 0.112, pediatric: 0.134, cumulative: 0.117) in the first year post-implant and then rises again in years of anticipated pump replacement. In nonimplant years, surgical revision rates remain low (adult: 0.060, pediatric: 0.073, cumulative: 0.063) relative to published rates and stable.

Safety can be further evaluated in this registry through the use of patient survival from therapy discontinuation—that is, the percentage of patients who remain on therapy at specific time points. Study exit (whether due to an adverse or device event or to inactive products) was considered registry discontinuation due to an unacceptable risk-benefit ratio. Therapy continued beyond 10 years for >75% of pediatric and adult patients in this registry, supporting the assertion of a positive risk-benefit determination, and high patient acceptance for ITB therapy.

Finally, these data offer the opportunity to further assess the risk-benefit ratio of ITB, provided that elective pump replacements are accepted as a surrogate for continued patient satisfaction. For patients maintaining ITB therapy for the lifetime of the implanted pump, pump replacement discussions can begin as much as one year before the required replacement surgery. During this time, the patient, their caregivers, and the managing physician can fully weigh the known and understood therapy risks against actual benefits received. In contrast to the theoretical risks and benefits associated with the first pump implant, implanted patients understand the daily risks and concerns associated with ITB therapy and the benefits derived. Patients also progress through other therapeutic options before ITB therapy, giving them insight into other therapies, and often into the consequences of no spasticity treatment. Analysis of elective pump replacement is possible in those patients who have both remained active in the registry and progressed to pump EOS—approximately 7 years. Although only a subset of the 1743 enrolled patients are available for analysis in this cohort (364 initial implants and 98 replacement implants), the significance of 426 patients with long-term ITB therapy (average 72 months) available for analysis cannot be understated. That 99% of these patients (98.9% of ITB naïve patients), following careful consideration, opt for therapy continuation indicates high patient satisfaction.

Death was a reason for discontinuation for 178 (15%) patients in this cohort with 169 reported as unrelated to the device, therapy or procedure and nine deaths of unknown or unreported relation to device/therapy. Two of these nine deaths involved pump pocket infections in immunocompromised patients, which are not classified as device related: one patient treated with ITB for traumatic brain injury-related spasticity experienced a pump pocket infection within 1 month of implantation that progressed to sepsis and resulted in death; the second patient, being treated for CP-related spasticity, experienced pump pocket erosion and an MSSA-positive infection 2 years post-implantation that resulted in death. A third patient death was associated with urinary tract infection and meningitis and reported as not device related, although system explant was undertaken. No literature providing mortality statistics specific to ITB therapy in patients with severe spasticity across all indications was identified, but research has shown that mortality in CP is the same with or without ITB treatment (25) and that infection risk is higher in the pediatric population compared to adults (26).

Registries such as the PSR are not without limitations. Physicians maintain their standard clinical practice and this lack of a uniform treatment limits the possibility of reproducing study results in other studies or within a clinical practice. Patient follow-up continues until a definitive therapy-related event occurs (e.g., therapy discontinuation or death), but patients may discontinue registry participation due to study site closure or to a transfer of care (e.g., pediatric patients aging up to adult management), limiting full follow-up. This study has more adult than pediatric spasticity patients, possibly skewing the cumulative results in favor of lower risk patients, whose bodies are no longer growing. The duration of this study (~15 years) additionally spans changes in ITB therapy best practices. These limitations are balanced by the fact that site selection and physician participation include various practice levels, including academic medical centers and private practice. In addition, high data quality is expected as implanted patients maintain close contact with the prescribing physician to continue/discontinue therapy. Finally, we have used acceptance of pump replacement as a surrogate for patient satisfaction with ITB, but patients may possibly opt for a pump replacement as a routine, or for fear of baclofen withdrawal reactions.

CONCLUSIONS

The PSR provides the largest adult and pediatric patient cohort to date with results on the long-term safety and device performance of ITB in the treatment of severe spasticity of cerebral and spinal origin. Both low AE-related therapy discontinuation rates and high acceptance of surgical intervention for therapy continuation were observed. Although other unmeasured factors may contribute to continuation and satisfaction of ITB therapy, the patient/caregiver willingness to continue ITB and accept surgical and other therapy risks was extremely high.

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Authorship Statements

Dr. Schiess wrote the first draft and participated in statistical analysis execution, review and critique. Dr. Eldabe, Dr. Plunkett, Dr. Konrad, and Dr. Molus participated in statistical analysis execution, review and critique, and in writing review and critique. R. Spencer participated in statistical analysis review and critique, and in writing review and critique. K. Stromberg participated in statistical analysis design and execution. T. Weaver participated in research conception, organization and execution, in statistical analysis review and critique, and in writing review and critique.

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