Intrathecal Drug Delivery Systems for Cancer Pain: An Analysis of a Prospective, Multicenter Product Surveillance Registry

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BACKGROUND: The safety and efficacy of intrathecal drug delivery systems (IDDSs) for the treatment of cancer-related pain have been demonstrated in randomized controlled clinical trials (RCTs). Despite positive evidence for this therapy, IDDS remains underutilized to treat cancer pain. Real-world registry data augment existing safety and effectiveness data and are presented here to broaden awareness of this therapeutic option, needed for adequate cancer-related pain treatment, and as a viable tool addressing concerns with systemic opioid use.

METHODS: This prospective, long-term, multicenter (United States, Western Europe, and Latin America) registry started in 2003 to monitor the performance of SynchroMed Infusion Systems. Patient-reported outcomes were added in 2013. Before data acquisition, all sites obtained Ethics Committee/Institutional Review Board approval and written patient consent. The study was registered (NCT01524276 at clinicaltrials.gov) before patients were enrolled. Patients who provided informed consent were enrolled in the registry at initial IDDS implant or replacement.

RESULTS: Through July 2017, 1403 patients with cancer pain were enrolled and implanted. The average (minimum/maximum) age of patients was 59 years (13/93 years), with 56.6% female. The most frequent cancer types were lung, breast, colon/rectal, pancreatic, and prostate. The majority of patients whose registry follow-up ended (87%; 1141/1311) were followed through death, with 4.3% (n = 57) exiting due to device explant or therapy discontinuation; the remaining 113 (8.6%) discontinued for reasons such as transfer of care, lost to follow-up, and site closure. Pain scores within the cohort of patients providing baseline and follow-up data improved significantly at 6 (P = .0007; n = 103) and 12 (P = .0026; n = 55) months compared to baseline, with EuroQol with 5 dimensions (EuroQol-5D) scores showing significant improvement at 6 months (P = .0016; n = 41). Infection requiring surgical intervention (IDDS explant, replacement, pocket revision, irrigation and debridement, etc) was reported in 3.2% of patients.

CONCLUSIONS: Adequate and improved pain control in patients with cancer, even in advanced stages, with concurrent quality of life maintenance is attainable. Results from this large-scale, multicenter, single-group cohort supplement existing RCT data that support IDDS as a safe and effective therapeutic option with a positive benefit–risk ratio in the treatment of cancer pain. (Anesth Analg 2020;130:289–97)

KEY POINTS

• Question: Do real-world registry data support the use of intrathecal drug delivery system (IDDS) as a treatment option for cancer-related pain?
• Findings: Overall device performance and safety along with positive outcomes in pain management and quality of life for the subset of patients available for analysis were demonstrated.
• Meaning: Real-world data and patient outcomes presented here add to the available evidence supporting the use of IDDS for cancer-related pain.

PLAIN LANGUAGE SUMMARY

Based on a study of 1403 patients, cancer-related pain can be successfully and safely treated with medication delivery by an implantable pump directly to the spinal cord.

GLOSSARY

5D = 5 dimension; 5L = 5 level; AE = adverse event; ASA = American Society of Anesthesiology; ASCO = American Society of Clinical Oncology; CI = confidence interval; CMM = conventional medical management; EQ-5D-5L = EuroQol with 5 dimensions (mobility, self-care, usual activities, pain/discomfort, and anxiety/depression) and 5 possible responses (having no problems, having slight problems, having moderate problems, having severe problems, and being unable to do/having extreme problems); EQ-5D-VAS = EuroQol with 5 dimensions Visual Analog Scale; EQ Health-VAS = EuroQol Health Visual Analog Scale (100: best; 0: worst); EQ-VAS = EuroQol Visual Analog Scale; FDA = Food and Drug Administration; IDDS = intrathecal drug delivery system; IQR = interquartile range; ISPR = Implantable Systems Performance Registry; MedDRA = Medical Dictionary for Regulatory Activities; MRI = magnetic resonance imaging; n = number (usually of subjects or occurrences); NPRS = numerical pain rating score (0: no pain; 10: worst pain); NCCN = National Comprehensive Cancer Network; PSR = Product Surveillance Registry; RCT = randomized controlled trial; SAE = serious adverse event; SD = standard deviation; STROBE = Strengthening the Reporting of Observational Studies in Epidemiology; VAS = visual analog scale; WHO = World Health Organization
The World Health Organization (WHO) cancer pain treatment guidelines\(^1\) identified inadequate cancer pain management as a global health concern, with the WHO analgesic ladder developed to support stepwise progression to strong opioids as necessary to adequately control cancer-related pain. The 2018 National Comprehensive Cancer Network (NCCN) pain guidelines recommend a similar stepwise algorithm, and link survival to disease/symptom control, including pain management, and to quality of life.\(^2\) Recent studies, however, indicate that cancer pain remains undertreated in 25\(^{\circ}\)\(^3\) to 77\(^{\circ}\)\(^4\) of patients (lack of adherence to WHO guidelines), with undertreatment rates unchanged over the past 20 years.\(^5\) In addition, the 5-year survival rate for all cancer types has increased to 65%,\(^6\) and many survivors experience chronic pain—with the American Society of Clinical Oncology (ASCO) policy indicating their need for long-term use of opioids.\(^7\) The current US opioid epidemic has increased scrutiny of systemic opioid use,\(^8\) so finding acceptable alternative treatments for refractory cancer pain and chronic pain is pressing.

Intrathecal drug delivery systems (IDDSs) administer Food and Drug Administration (FDA)-approved preservative-free morphine sulfate or ziconotide directly to the spinal cord. IDDS has demonstrated improved cancer pain management compared to conventional medical management (CMM) and placebo\(^10\)–\(^12\) and in retrospective and observational studies\(^13\)–\(^14\) for patients unresponsive to escalating systemic opioid doses or experiencing intolerable side effects. For those patients, IDDS facilitates systemic medication reduction or elimination with associated risk reduction\(^15\)–\(^15\) and significant cost savings.\(^16\) This single-group cohort study presents a compilation of 14 years of observational IDDS data, prospectively collected and monitored for compliance, on 1403 registry participants with cancer pain.

**METHODS**

This study followed appropriate guidelines for a cohort study (Strengthening the Reporting of Observational Studies in Epidemiology [STROBE] checklist). Before data acquisition, all sites obtained Ethics Committee/Institutional Review Board approval and patient consents for all subjects. The study was registered before patients were enrolled (NCT01524276; responsible party: Medtronic; date of registration: January 2012).

**Registry Description**

The Implantable Systems Performance Registry (ISPR; ClinicalTrials.gov Identifier: NCT01524276), initiated in 2003, is described in detail by Konrad et al.\(^17\) Results presented here include data collected on patients implanted with the SynchroMed II Infusion System (Medtronic, Inc, Minneapolis, MN) and enrolled in the ISPR (2003–2012) as well as those implanted and enrolled in the 2013 amended registry, referred to as the “Product Surveillance Registry (PSR),” which is ongoing and collectively referred to as the “registry.” The registry platform was designed to conduct ongoing nonrandomized, active prospective postmarket surveillance under a common protocol with specific appendices for neuromodulation products/therapies, enrolling patients with eligible, commercially available products. Product performance and patient outcomes are assessed compared to baseline, but no comparison group is included. Data collection aligns with routine clinical practice and was, therefore, not limited to on-label drug administration. Registry sites contributing to these data are noted in the Acknowledgments.

**Patients**

Potential registry patients are identified from the practices of participating physicians as meeting specific indications (eg, Chronic Intractable Malignant Pain) for the SynchroMed II Infusion System and are enrolled at initial implant or at the time of replacement for a previously implanted pump. The patient or legally authorized representative provides written authorization and/or consent per institution and geographical requirements before 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 are ineligible. Data are only included for patients who consent to enroll. After enrollment, patients are followed per standard of care, with status updates obtained every 6 months through therapy discontinuation or registry exit.

Data Collection
Registry evolution has resulted in data collection changes over time. A numerical pain rating scale (NPRS) assessment was initiated in 2010, with a single pain score (0: no pain; 10: worst pain, assessed as “current pain”) collected during scheduled study visits. The EuroQol with 5 dimensions (mobility, self-care, usual activities, pain/discomfort, and anxiety/depression) and 5 possible responses (having no problems, having slight problems, having moderate problems, having severe problems, and being unable to do/having extreme problems) (EQ-5D-5L) and EQ Health Visual Analog Scale (EQ Health-VAS) assessments of quality of life were added to the registry in 2013. The United Kingdom value set18 was used to calculate the EQ-5D Index value, which ranges from 1 (best health state) to −0.285 (worst health state). A positive mean index value change indicates an improvement (ie, increase) in health. The EuroQol Visual Analog Scale (EQ-VAS) assesses patient-reported health on that day, ranging from 0 (worst health) to 100 (best health), and a positive mean visual analog scale (VAS) change indicates an improvement in patient-reported health. Neither of these patient-reported outcomes allowed for retrospective collection of data. Thus, limited baseline pain and EQ-5D scores were available for older registry entries.

The American Society of Anesthesiology (ASA) physical status scores were collected as standard practice at the time of IDDS implant consideration at a single center on a subset (n = 649) of enrolled subjects.

Safety data currently collected in the registry include all events that appear or worsen after enrollment and are a result of implanted or external components of the IDDS, the implant procedure, or the infusion therapy. Adverse events (AEs) were categorized using Medical Dictionary for Regulatory Activities (MedDRA) criteria and terminology.

Analytic Methods
Data included in this analysis were collected through July 31, 2017, from patients in the registry who were treated with IDDS for cancer pain. Summary statistics are presented either as percentages for categorical variables or mean (standard deviation [SD]; or minimum/maximum) or median (interquartile range [IQR]) for continuous values.

Patient survival was defined as freedom from all-cause mortality and estimated using Kaplan–Meier survival analysis methods. Survival time was defined as months from the patient’s first implant recorded in the registry to death. Patients who remained alive were censored at their last follow-up in the registry.

Pain and EQ-5D patient-reported outcomes were analyzed for therapy-naïve patients who were enrolled in the registry with their first pump implant and provided pain/EQ-5D baseline data before pump implant. Analysis is provided on 2 cohorts of patients: those with 6-month data collected and those with 12-month data collected; all patients with required data for analysis were included in each analysis set. Paired t tests were used to test within-patient change from baseline to follow-up for paired data. The Hochberg method19 was used to adjust the significance level for the multiple statistical tests that were performed (change in pain, EQ-5D, and EQ-VAS from baseline to 6 and 12 months).

Events related to product performance, requiring surgical intervention (including infections requiring surgical intervention), and reports of patient death have been collected consistently since registry inception. Summaries of these events include all patients. AE reporting was expanded in 2010 to include serious AEs (SAEs) of any etiology related to the device. The subset of active and newly enrolled patients after this protocol change is included in the analysis of AEs. Safety summaries are presented as the percentage of patients who experienced ≥1 event, with 95% Wilson score confidence intervals (CIs).

RESULTS
Patient Demographics and Implant Details
A total of 7867 IDDS patients were enrolled in the registry at 64 sites across the United States, Europe, and Latin America, with 1403 from 37 sites being treated for cancer pain (Supplemental Digital Content, Figure 1, http://links.lww.com/AA/C934). The majority of patients were enrolled at 1 registry site (n = 1136, 81.0%, Phoenix, AZ), with an additional 7 sites contributing 12.7% of patients (n = 178) and the remaining 29 sites each contributing ≤10 patients.

The average (minimum/maximum) age of the patients was 59 years (13/93 years), with 56.6% female (Table 1). The SynchroMed II 40-mL pump was the prevalent pump implanted (1267/1505 pumps, 84.2%), and the InDura Model 8709 catheter the most common catheter (980/1535 catheters, 63.8%), with Ascenda Models 8780 and 8781 (53 and 282 catheters, respectively; total 335 of 535 catheters, 21.8%) also implanted. Medical history data indicated median (IQR) duration from cancer diagnosis to implant of 28.9 months (14.9–66.0 months), with median (IQR) duration from implant to death/last visit of 3.2 months (1.2–9.4 months).

ASA physical status for 91.1% (n = 591) of the analyzed subgroup was III or IV, with 8.9% (n = 58) ASA
with an existing cancer diagnosis, there were no ASA I subjects. The average ASA score within this cohort subset was 3.12.

**Table 1. Demographics and Baseline Data of Patients With Cancer Pain Enrolled in the Product Surveillance Registry**

| Category                                      | Value                  |
|-----------------------------------------------|------------------------|
| Age at enrollment, y                         | 59 (13/93)             |
| Female, %                                     | 56.6                   |
| Male, %                                       | 43.4                   |
| Type of cancer, n (%)                         | 592*                   |
| Lung                                          | 89 (15.0)              |
| Breast                                        | 65 (11.0)              |
| Colon/rectal                                  | 64 (10.8)              |
| Pancreatic                                    | 49 (8.3)               |
| Prostate                                      | 35 (5.9)               |
| Bladder                                       | 23 (3.9)               |
| Other/unknown                                 | 267 (45.1)             |
| Months from diagnosis to implant              | 28.9 (14.9–66.0)       |
| Median (IQR range)                            | 3.2 (1.2–9.4)          |
| Baseline pain score                           | 6.8 (2.4)              |
| Mean (SD)                                     | 6.8 (2.4)              |
| Median EQ-5D Index score                      | 0.372 (0.269)          |
| Mean (SD)                                     | 0.379                  |
| ASA physical status at enrollmentb            | 3.12                   |

**Patient Survival**

Patient postimplant survival was 39%, 24%, 16%, 11%, and 5% at 0.5, 1, 2, 3, and 10 years, respectively (Figure 1). As of July 31, 2017, 92 IDDS patients remained actively enrolled in the registry and 1311 had exited the registry. Complete follow-up (not lost to follow-up or withdrawal) was high, with the majority of patients (1141/1311; 87%) followed from implant through death, and only 4.3% exiting the study due to device explant or therapy discontinuation; the remaining 113 patients (8.6%) discontinued for reasons such as transfer of care, lost to follow-up, and site closure. Among patients exiting the registry due to death, >90% (1052/1141) expired due to disease-related (neoplasm) causes. Only 2 deaths were possibly associated with therapy: one was reported as infection with death secondary to postoperative pneumonia after device implantation, and one was due to pulmonary embolus secondary to drug withdrawal as a result of missed pump refill. Of the 87 remaining patient deaths, none was associated with product performance, therapy, or surgical implantation. The duration from implant to last patient follow-up ranged from <1 month to >14 years.

**Pain**

Data were analyzed on the subcohort of 283 patients for whom baseline pain scores were available; these patients reported an average baseline pain score of 6.8 (SD, 2.4). A subset of these 283 patients had both baseline and follow-up (6 and/or 12 months) pain

![Figure 1. Patient survival from all-cause death through (A) 12 mo and (B) 126 mo. Shaded area represents the 95% confidence interval at that time point. Number of patients at risk are shown at select months of follow-up. Data are shown when there are ≥20 patients in each 3-month interval.](image-url)
scores. Under an adjusted significance level of .017, these patients demonstrated statistically significant improvement in average pain from baseline (6.6; SD, 2.4) to 6 months (5.5; SD, 2.6, n = 103), with an average change of −1.1 (95% CI, −0.5 to −1.7; P = .0007), and from baseline (6.9; SD, 2.3) to 12 months (5.4; SD, 2.5; n = 55), with an average change of −1.4 (95% CI, −0.5 to −2.3; P = .0026) (Figure 2).

Quality of Life

Patient-reported quality of life, as indicated by the EQ-5D Index value and the EQ-5D Health-VAS, demonstrated statistically significant improvement compared to baseline at 6 months (n = 41) under an adjusted significance level of .017. The average EQ-5D Index value improved from 0.386 (SD, 0.252) to 0.556 (SD, 0.252) with an average change of +0.171 (95% CI, 0.069–0.273; P = .0016). The average EQ-5D Health-VAS score improved from 45.2 (SD, 21.6) to 58.2 (SD, 20.7) with an average change of +13.0 (95% CI, 4.5–21.5; P = .0036). Change in quality of life from baseline to 12 months, however, was not statistically significant (P = .13, adjusted significance level of .025, for EQ-5D Index value; and P = .22, adjusted significance level of .05, for VAS) (Figure 3).

Safety

Within the full cancer pain cohort (n = 1403), infection requiring surgical intervention (IDDS explant, replacement, pocket revision, irrigation and debridement, etc.) was reported in 3.2% (95% CI, 2.4–4.3) of patients. Pneumonia was the only reported infection resulting in death (n = 1), and possibly attributed to the device or a therapy-related surgical procedure. Events defined as product performance related (events with an etiology associated with pump, catheter, or patient programmer performance) and occurring in >1% of patients were catheter dislodgement (3.8%; 95% CI, 2.9–4.9), pump motor stall (1.8%; 95% CI, 1.2–2.6), catheter occlusion (1.5%; 95% CI, 0.9–2.3), catheter kink (1.5%; 95% CI, 0.9–2.3), and catheter break/cut (1.2%; 95% CI, 0.8–1.9). Magnetic resonance imaging (MRI) exposure, a potential cause of motor stall, has been actively tracked within the registry since 2013. A total of 73 MRIs was reported for 51 patients, with all but 1 reported as event-free. All the MRI-induced motor stalls recovered within the expected 24-hour period.

Of the 706 patients who were followed after the 2010 expanded AE data collection, 40% (279/706; 95% CI, 36–43) experienced ≥1 AE that was related to the device components, implant procedure, or delivery of therapy. The most frequently occurring AEs were adverse drug reaction (24.5%; 95% CI, 21.5–27.8) and medical device site pain (10.1%; 95% CI, 8.1–12.5). Sixty-eight SAEs were reported in 54 patients (7.65%) (Table 2).

DISCUSSION

Data from this large-scale multicenter registry supplement existing randomized controlled trial (RCT) data demonstrating the safety and effectiveness of IDDS as a therapeutic option for the treatment of cancer pain. Both pain and quality of life scores demonstrated significant improvement from baseline at 6 months after IDDS implantation, with significant improvement from baseline in pain scores at 12 months. Only 4.3% of discontinued patients exited the study due to device explant or therapy discontinuation. Infection requiring surgical intervention occurred in 3.2% of patients.

The WHO1 and NCCN2 guidelines sought to address undertreatment of pain by optimizing medication selection and escalating dose to maintain adequate pain control with disease progression. Systemic opioids represented the standard of care in both

Figure 2. Patient-reported outcomes: pain. Average pain scores and 95% confidence intervals for patients with paired baseline and 6- or 12-mo assessment data. Pain scores range from 0 (no pain) to 10 (worst pain).
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Table 2. Therapy-Related Serious Adverse Events

| MedDRA SOC | MedDRA Preferred Term | No. of Serious Events | No. (%) of Patients n = 706 | 95% Confidence Interval |
|------------|-----------------------|-----------------------|----------------------------|-------------------------|
| Infections and infestations | Subtotal | 17 | 17 (2.41) | 1.51–3.82 |
| | Implant site infection | 10 | 10 (1.42) | 0.77–2.59 |
| | Medical device site infection | 3 | 3 (0.42) | 0.14–1.24 |
| | Wound infection | 2 | 2 (0.28) | 0.08–1.03 |
| | Incision site infection | 1 | 1 (0.14) | 0.03–0.80 |
| | Meningitis | 1 | 1 (0.14) | 0.03–0.80 |
| Psychiatric disorders | Subtotal | 5 | 4 (0.57) | 0.22–1.45 |
| | Withdrawal syndrome | 5 | 4 (0.57) | 0.22–1.45 |
| Nervous system disorders | Subtotal | 9 | 9 (1.27) | 0.67–2.40 |
| | Cerebrospinal fluid leakage | 5 | 5 (0.71) | 0.30–1.65 |
| | Headache | 2 | 2 (0.28) | 0.08–1.03 |
| | Hypoesthesia | 1 | 1 (0.14) | 0.03–0.80 |
| | Spinal cord hematoma | 1 | 1 (0.14) | 0.03–0.80 |
| Respiratory, thoracic, and mediastinal disorders | Subtotal | 2 | 2 (0.28) | 0.08–1.03 |
| | Acute respiratory failure | 1 | 1 (0.14) | 0.03–0.80 |
| | Respiratory failure | 1 | 1 (0.14) | 0.03–0.80 |
| General disorders and administration site conditions | Subtotal | 29 | 23 (3.26) | 2.18–4.84 |
| | Adverse drug reaction | 12 | 10 (1.42) | 0.77–2.59 |
| | Pain | 9 | 8 (1.13) | 0.58–2.22 |
| | Drug withdrawal syndrome | 7 | 5 (0.71) | 0.30–1.65 |
| | Medical device site hematoma | 1 | 1 (0.14) | 0.03–0.80 |
| Injury, poisoning and procedural complications | Subtotal | 6 | 6 (0.85) | 0.39–1.84 |
| | Overdose | 3 | 3 (0.42) | 0.14–1.24 |
| | Postlumbar puncture syndrome | 1 | 1 (0.14) | 0.03–0.80 |
| | Procedural pain | 1 | 1 (0.14) | 0.03–0.80 |
| | Wound dehiscence | 1 | 1 (0.14) | 0.03–0.80 |
| Total | | 68 | 54 (7.65) | 5.91–9.85 |

Abbreviations: AE, adverse event; MedDRA, Medical Dictionary for Regulatory Activities; SOC, System Organ Class.

*https://www.meddra.org/.

**AE seriousness was added to the registry in April 2010. This table includes AEs categorized as serious in the active patients (n = 706) from April 2010 to July 31, 2017.

*The total number of patients with events may not represent the sum of all rows because a patient may have experienced >1 type of event.

Figure 3. Patient-reported outcomes: quality of life. A, Average EQ-5D Index scores and 95% confidence intervals for patients with paired baseline and 6- or 12-mo assessment data. EQ-5D Index scores range from −0.285 (worst health state) to 1 (best health state). B, Average EQ-5D Health-VAS and 95% confidence intervals for patients with paired baseline and 6- or 12-mo assessment data. EQ-5D Health-VAS scores range from 0 (worst health) to 100 (best health). EQ-5D indicates EuroQol with 5 dimensions; EQ-5D Health-VAS, EuroQol Health Visual Analog Scale.
guidelines. The NCCN guidelines were expanded to include palliative and interventional referrals but do not address the lack of effectiveness data for long-term opioid use and inherent risks. Despite accepted use, there are no existing RCT data demonstrating long-term effectiveness of systemic opioids, and they are also associated with tolerance, hyperalgesia, addiction, sleep disorders, breathing/brain deregulation with early dementia, hypothalamic–pituitary deregulation, fractures, depression, and side effects including lethargy, sedation, nausea with vomiting, mental cloudiness, and constipation that significantly impact quality of life. Treatment options for cancer pain should also consider potential patient survival by addressing both immediate and longer-term pain management needs. Two prospective multicenter studies (IDDS + CMM versus CMM and active treatment versus placebo) provide RCT support for IDDS effectiveness in pain relief, toxicity reduction, and quality of life improvement. Compared to CMM, IDDS-treated patients have also had lower utilization and total medical costs in the first year after implantation, driven by the savings in hospitalization and emergency department visits. A recent retrospective claims analysis found savings of $15,142 ($P = .0097) at 2 months and $63,498 ($P = .0097) at 12 months compared to CMM after starting IDDS.

Perceived IDDS risks associated with implantation and management have been identified as a limiting factor in therapy acceptance, and a possible reason for the delayed referral to pain physicians for treatment after cancer diagnosis. IDDS risk data presented here, however, compare favorably to that for intravenous ports placed for medication delivery. ASA scores collected on a subset of subjects additionally indicate the health status of a large proportion of patients included in this analysis were predisposed to increased surgical risk. Surgical-related SAEs included infection (2.41%), postdural puncture headaches/cerebral spinal fluid leaks (1.27%), and pump pocket hematoma (0.28%). IDDS implantation, as a minimally invasive surgery, resulted in clinically significant improvements to pain, function, and quality of life.

Therapy-related SAEs included respiratory failure (0.14%), adverse drug reactions (1.42%), pain (1.13%), and overdose (0.42%), none of which resulted in death. Most of these events were likely secondary to opioid rotation, drug delivery escalation, or return of underlying symptoms. Drug withdrawal SAEs (0.71%) were related to missed pump refill appointments, catheter complications, dosing changes, or pump motor stall associated with off-label medication usage. SynchroMed II motor stall risks were addressed by a redesign in collaboration with the FDA addressing gear corrosion. The SynchroMed II pump motor may stall during MRI, but the pump should resume normal function after MRI exposure. No permanent motor stalls after MRI were reported in this cohort of patients; temporary motor stalls were reported but resolved after MRI, with no reports of post-MRI drug withdrawal or sequela.

Analysis of pain and quality of life measurements within the subset of patients providing both baseline and follow-up data offer additional, albeit noncomparative, evidence that IDDS is an effective treatment option for patients experiencing significant cancer pain. Compared with baseline, patients followed through 6 months had improved pain and functional status, with those followed up to 1 year maintaining improved pain management. Patient survival data indicate that a subset of patients have survived through 10 years of active treatment, which necessitates ≥1 pump replacement. Given the increasing survival and incidence of chronic cancer pain in long-term survivors, IDDS therapy offers a long-term treatment option that avoids the toxicities associated with oral medications.

Uncontrollable pain is often cited as the most feared aspect associated with a cancer diagnosis, with severity of pain associated with decreased treatment compliance, reduced survival, and increased rates of suicidal ideation and suicide. In this cohort of patients, baseline pain was classified as severe, with a baseline median pain score in the presence of standard pain control measures of 7/10. Although not directly assessed in this registry, earlier intervention with IDDS with improved pain control and reduction in side effects associated with oral medications may positively impact overall survival. Patients here had a median time between cancer diagnosis and IDDS therapy of ≥2 years. Although this may represent a delay in onset of severe pain in some of these patients, it likely represents a prolonged duration of poorly controlled pain in many, emphasizing the importance of a pain management plan, established at cancer diagnosis in coordination with pain specialists, addressing timely advancement from conservative therapies to advanced interventions when appropriate. Inclusion of these specific pain management plans as part of palliative care in Survivorship Care Plans, with buy-in from the diverse providers seen by cancer patients, would serve well in achieving the goals established by the WHO Guidelines >20 years ago.

Opioids remain the mainstay of treatment for cancer pain in the United States, but their identified risks and the opioid epidemic have placed undue burden on the prescribing provider and may result in risk of undermanagement of pain. The safety and outcome data presented here, as well as reduced risk of drug
diversion or medication surplus after patient demise, strongly support a more widespread acceptance of IDDS for cancer pain. Although IDDS is not free from all medication-related risks, consensus statements regarding the safety, side effects, and best clinical practices for IDDS,26–28 published regularly since 2000 for nonmalignant pain, offer additional support for wider acceptance of this needed therapy.

Limitations
The registry data were collected using protocols that allowed clinicians to maintain their standard clinical practice. Patient-reported outcomes were only added in the past few years, so data were presented for the more recent, limited subset of the registry cohort. AE collection expanded in 2010, but surgical interventions and reasons for discontinuation have been captured consistently since registry inception. Although large registries are becoming more widely accepted in the assessment of safety and patient outcomes, the registry is limited by not having a direct comparator. Only implanted patients continue and provide data in the registry; no concurrent nonimplanted group is available for comparison to those patients receiving IDDS therapy. In addition, most patients presented here were treated at a single center in the United States. This center demonstrates a higher referral pattern from oncology with more extensive experience with aggressive treatment of cancer-related pain. Comparison of patient outcomes from this center to that of all other centers indicated no difference in the rate of SAEs but did demonstrate a difference in AE rates (higher AE rate, statistically significant) as well as a difference in EuroQol Visual Analog Scale (EQ-5D-VAS) scores at 6 months (greater improvement, statistically significant) for this center. These data likely reflect more aggressive dosing and titration patterns of IDDS therapy. Predictably, the most frequently occurring AEs were medication related, as well as medical device site pain. Finally, the registry represents a heterogeneous patient population. Although all had cancer pain as the primary indication for IDDS, essentially all types of cancer were included, as were many different pain etiologies (ie, pain due to tumors or to cancer treatments), and patients were enrolled at various stages of disease.

CONCLUSIONS
This registry remains unique in terms of enrollment numbers and duration of follow-up for patients with cancer pain. Adequate and improved pain control in patients with cancer, even in advanced stages, with concurrent quality of life maintenance are attainable. Results from this large-scale, multicenter registry supplement existing RCT data that support IDDS as a safe and effective therapeutic option with a positive benefit–risk ratio in the treatment of cancer pain.

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Conflicts of Interest: L. M. Stearns is a paid consultant for Medtronic, Flowonix, Nevro, and Boston Scientific and receives research support from Medallion Therapeutics, Inc, Medtronic, Boston Scientific, and Piramal for services unrelated to the current research.
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