Neuromodulation with electrical field stimulation of dorsal root ganglion in various pain syndromes: a systematic review with focus on participant selection

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Objective: We conducted a systematic review about patient selection, efficacy, and safety of neuromodulation with electrical field stimulation (EFS) of dorsal root ganglion (DRG) in various painful conditions. We also analyzed conclusion statements as well as conflict of interest and financing of the included studies.

Methods: All study designs were eligible for inclusion. We searched MEDLINE, CINAHL, Embase, PsycINFO, and clinical trial registries until September 7, 2018. We assessed risk of bias by using Cochrane tool for randomized controlled trials (RCTs).

Results: Among the 29 included studies, only one was RCT, majority being case series and case reports. The evidence is based on studies with small number of participants (median: 6, range 1–152) with various painful conditions. Neuromodulation with EFS of DRG was mostly performed in participants who have failed other treatment modalities. Most of the authors of the included studies reported positive, but inconclusive, evidence regarding efficacy of neuromodulation with EFS of DRG. Meta-analysis was not possible since only one RCT was included.

Conclusion: Available evidence suggest that neuromodulation with EFS of DRG may help highly selected participants with various pain syndromes, who have failed to achieve adequate pain relief with other pharmacological and nonpharmacological interventions. However, these findings should be confirmed in high-quality RCTs with sufficient numbers of participants.

Keywords: DRG, pain, neuropathic pain, chronic pain, neurostimulation, electrical stimulation

Introduction

Neurostimulation is a widely used therapeutic approach to treat various painful conditions including complex regional pain syndrome (CRPS),¹–³ chronic low back pain (LBP),⁴–⁶ groin pain,⁷,⁸ and pelvic pain.⁹,¹⁰ Neurostimulation as a therapeutic method uses electrical energy, that is, electrical field stimulation (EFS) in order to functionally activate or inhibit neuronal groups, networks, or pathways and to achieve pain relief.¹¹,¹²

Primary sensory neurons and their somata in dorsal root ganglia (DRGs) are important sites where pathologic changes that lead to neuropathic pain occur, creating an opportunity for selective neuromodulation.¹³,¹⁴ Data from animal model studies demonstrated that neuromodulation with EFS of DRG has several advantages compared to spinal cord stimulation (SCS), allowing more precise positioning of stimulation leads with increased flexibility and reduced contact size and spacing, leading to better pain relief.¹⁵ Although the neuromodulation mechanism of DRG EFS has
not been clearly elucidated, several animal studies showed that the branching point into peripheral and central process of pseudounipolar sensory neuron, that is, T-junction,\(^1\) has a filtering role in the propagation of action potentials from periphery to the spinal cord\(^1\) and can be used as a target for therapeutic stimulation that can lead to reduction of pain.\(^1\)\(^5\)\(^1\)\(^7\)

Our group has recently published a systematic review about the use of neuromodulation in the context of pain from in vivo and in vitro preclinical animal model studies that showed that neuromodulation with EFS of DRGs had generally positive therapeutic effects in the context of pain.\(^1\)\(^9\) However, we found low methodological quality of included studies, as well a need for using standardized models and outcomes to better understand how DRG stimulation reduces pain in animal models.\(^1\)\(^9\) Heterogeneity of preclinical models used to study neuromodulation in the context of pain precludes any quantitative synthesis of results from different studies.\(^1\)\(^9\)

Despite scarcity of data from preclinical models, DRG stimulation has already been used extensively in clinical settings.\(^1\)\(^2\) Moreover, in February 2016, the US Food and Drug Administration granted premarket approval to Axium Neurostimulator System (Spinal Modulation, Inc., Menlo Park, CA, USA, recently bought by Abbott Laboratories, Sunnyvale, CA, USA) after demonstration of its beneficial effect based on the ACCURATE study of Deer et al.\(^1\) The ACCURATE study was the only randomized controlled trial (RCT) performed in the field comparing neuromodulation with EFS of DRG with SCS for the treatment of CRPS and causalgia, with 152 participants. The study results demonstrated higher treatment success rate of DRG EFS neuromodulation compared to SCS.\(^1\) CRPS, for which Axium Neurostimulator System was approved, is defined as chronic pain of neuropathic origin after injury of limbs such as fractures, surgery or sprains, limb immobilization or as a reflection of internal neural damage.\(^2\)\(^0\) Reported prevalence of CRPS is <2\% in most retrospective series.\(^2\)\(^1\)

Several reviews about neurostimulation of DRG have been published recently. However, they mostly had a narrow focus, limited to painful condition or neuromodulation target, and they all had a number of methodological limitations.\(^2\)\(^2\)\(^2\)\(^3\) Recently published best practices on DRG stimulation by Neuromodulation Appropriateness Consensus Committee (NACC) gave a comprehensive overview of the topic, with focus on stimulation devices and procedure techniques, whereas selection of participants was mentioned briefly.\(^2\)\(^4\) The aim of this systematic review was to create comprehensive evidence synthesis about efficacy and safety of neuromodulation with EFS of DRG for the treatment of various painful conditions, with particular emphasis on participant selection.

**Methods**

**Study design**

We conducted a systematic review in accordance with the methods and guidelines from the Center for Reviews and Dissemination (CRD)\(^2\)\(^5\) and the PRISMA statement.\(^2\)\(^6\)

**Protocol and registration**

The protocol of this systematic review was developed a priori and registered in the PROSPERO database (registration number: CRD42017076502).

**Eligibility criteria**

**Participants**

We included primary studies that analyzed participants with any type of pain syndrome and any intensity of pain.

**Interventions**

EFS of DRG, regardless of the parameters of stimulation.

**Comparators**

Any type of comparator was eligible. We also included studies that had analyzed EFS neuromodulation of DRG without a comparator group.

**Outcomes**

The main outcome measures were pain intensity and serious adverse events (SAEs) as they were defined in included manuscripts. Secondary outcome measures were any other safety data and any other pain-related outcomes. We considered all follow-up periods with no cutoff criteria.

**Study designs**

All study designs were eligible, including case reports. Although RCTs are considered the highest level of evidence of interventions in medicine, we were concerned that few RCTs were conducted in this field and that exclusion of nonrandomized study designs (NRSDs) would give an incomplete summary of the current evidence-base about the effects of the analyzed intervention in terms of efficacy and safety. We used the Cochrane Handbook definitions to determine the study design if the study design was not explicitly described in the manuscript. If a study reported cases, we considered it to be case series if it presented >10 participants, according to the definition of the Cochrane Handbook. We also reported study design definitions given...
by authors of included studies to present heterogeneity among definitions of NRSDs.

Information sources
We searched the following four databases: MEDLINE, CINAHL, Embase, and PsycINFO. We also searched ClinicalTrials.gov and WHO’s International Clinical Trial Registry Platform to identify the ongoing studies. Databases were searched from the date of their respective inceptions and the date of the last database search was September 7, 2018, whereas for trials registry, the date of last search was October 2, 2018.

References and citations of the included studies and any potentially relevant reviews were analyzed in order to find additional eligible studies that may not have been retrieved by the database search.

Search strategy
A computer-based search strategy was designed and conducted with the assistance of an expert medical librarian, who also peer-reviewed the final version of the search strategy. Search strategy for MEDLINE via OVID (Table S1) was developed first and adapted for other databases subsequently. Studies published in any language were considered. Searches were conducted separately in each database, and subsequently the records were exported to EndNote X5 citation software (Clarivate Analytics, Boston, MA, USA). Duplicates were removed, first by software and then manually. Reference lists of all the included studies were searched. Citations and references of the included studies were downloaded from Web of Science.

Study selection
Two authors independently screened titles and abstracts of the bibliographic records retrieved via the database search (SD and LFH). Two authors also independently screened full-text manuscripts of potentially relevant studies (IV and TM). In each step, disagreements were resolved via discussion or involvement of a third author (DS). When involvement of a third author was deemed necessary, the third author would suggest the solution, with arguments, and this was in all cases accepted by the co-authors.

Data collection process and data items
A data collection form was developed for this study and piloted using five included studies and subsequently revised the form as appropriate. Two authors (IV and TM) independently extracted data in duplicate. The following data were extracted: name of the first author, year of publication, study design, intervention and comparator, number of participants, participant characteristics including inclusion and exclusion criteria, baseline characteristics, previous therapy and painful condition, follow-up period, parameters of stimulation and stimulator used, position of leads, studied outcomes, and study results regarding efficacy and safety of intervention.

Risk of bias (RoB) assessment
To assess RoB in the included studies, we used the Cochrane RoB tool for RCTs, which has seven domains, addressing bias related to random sequence generation, allocation concealment, blinding of participants and personnel, blinding of outcome assessors, incomplete outcome data, selective reporting, and other potential sources of bias. For each domain, we reported our judgment, that is, whether the risk was low, high, or unclear, and we provided a supporting comment, which explained our judgment. RoB was assessed per domain level; we did not assess RoB on outcome level, and we did not assess overall RoB on an entire study level. RoB assessment was included in our narrative analysis and conclusions.

We aimed to use the Risk of Bias in Non-randomized Studies of Interventions (ROBINS-I) tool for cohort studies, but we did not have such studies in our sample. Two authors independently analyzed RoB (KV and SD), and discrepancies were resolved by the third author (LP).

Synthesis of results
We grouped the results according to painful condition treated with DRG stimulation. Studies that included participants with multiple etiologies were grouped into the painful condition category with highest number of participants and mentioned in other categories if relevant. The results are presented in a narrative and tabular form. We planned to conduct meta-analysis of outcomes from RCTs, but meta-analysis was not possible since only one RCT was included. In addition, we analyzed reporting of conclusion statements for efficacy and safety in manuscript abstracts. We extracted verbatim those conclusion statements and divided them into five categories: positive conclusive, positive inconclusive, negative conclusive, negative inconclusive, and not reported. We categorized as inconclusive conclusion statements that used conditional wording about the efficacy or safety and/or indicated that more evidence is needed.

Results
Electronic databases searches yielded 2,811 records. An additional 1,473 studies were identified through supplementary search of references, citations, and other reviews. After identifying and removing duplicate records, 2,133 unique
records remained for eligibility determination and inclusion. We analyzed a total of 39 manuscripts in full text, excluded 10 of them, and finally included 29 studies in narrative synthesis. A flowchart is presented in Figure 1, whereas characteristics of included studies are described in Tables 1–3. Studies excluded from further qualitative synthesis (N=10) and reasons for their exclusion are presented in Table S2. Ongoing studies found in clinical trial registries are presented in Table S3.

Included studies had various study designs: 1 RCT, 8 before and after comparisons, 2 case series, and 18 case reports. Liem et al (2013, 2015) reported the results of the same study with data shown for 6 months3 and 12 months follow-up time periods; 2 so we considered both of them. Two studies from van Velsen et al analyzed the same patient; therefore, we left both the references but included it in analysis only once.29,30 Due to a large number of case reports and case series, overall, the included studies were with very small median number of participants 6 (range: 1–152). Only one included RCT and two observational studies had higher number of participants.

Pain syndromes analyzed in included studies

Included studies analyzed the following painful conditions: CRPS,1–3,31–36 LBP,5,6,37–40 groin pain,7,41 pelvic girdle pain,9,10 peripheral neuropathy,29,30,42 peripheral diabetic neuropathy,43 phantom limb pain,44 chronic intractable pain in the coccyx,45 chronic testicular pain,46 anterior cutaneous nerve entrapment syndrome (ACNES),47 loin pain hematuria syndrome (LPHS).48

Studies awaiting classification

No results have yet been published for several completed clinical trials retrieved via search of clinical trials registries. One study was classified as completed with no results (NCT02169401). The trial authors informed us that the manuscript has been submitted. Other studies were classi-
Table 1 Characteristics of studies about EFS of DRG

| Study design | Number of participants | Follow-up | Outcome measures | Results: efficacy for pain intensity | Results: SAE | Results: any other safety data |
|--------------|------------------------|-----------|-----------------|-------------------------------------|--------------|--------------------------------|
| Deer et al (2017)¹ | CRPS and causalgia (32 participants) | DRG stimulation: 76 SCS: 76 | 3, 6, and 12 months | - Pain intensity by VAS; treatment success rate composed of i) >50% VAS pain relief after trial phase, ii) VAS score at 3 months reduced from baseline by >50%, and iii) did not experience stimulation-related neurological deficit • Positional effect on paresthesia intensity • Quality of life by SF-36 • Mood by POMS • Pain severity by BPI • Subject satisfaction • Stimulation specificity • VAS percentage change • AEs | DRG group: Baseline VAS: 80.6 VAS at 3 months: 13.1 VAS at 12 months: 15.0 SCS group: Baseline VAS: 80.7 VAS at 3 months: 23.8 VAS at 12 months: 26.5 The proportion of subjects who achieved treatment success at 3 months in DRG arm (81.2%) is statistically greater than in SCS arm (55.7%) At 12 months success was in DRG arm (74.2%) and 53% in SCS arm | 21 SAEs occurred in 19 subjects, 8 in DRG arm and 11 in SCS arm. The rates of SAE were 10.5% in DRG arm and 14.5% in SCS | 52 procedure-related AEs by 35 subjects in DRG arm and 29 AEs by 20 subjects in SCS arm |
| Liem et al (2013)² | CRPS (9 participants) and FBSS (8 participants) | Trial period: 59 Permanent implantation: 32 | After TNS and 1 week, 2, 3, and 6 months after INS | - AE rate and paresthesia generation • Pain intensity by VAS • Quality of life by EQ-SD • Mood by POMS • Physical functioning by BPI | Baseline VAS: 77.6±2.1 VAS after TNS: 26.1±3.4, reduction by 66.1% VAS 1 week after INS: 34.9±4.3, reduction by 55.1±5.5% VAS at 2 months (N=22): 39.5±6.6, reduction by 50.7±8% VAS at 3 months (N=30): 38.4±5.7, reduction by 50.8±7% VAS at 6 months (N=25): 33.5±6.0, reduction by 56.3% | 9 SAEs in 24 subjects | 61 AEs reported |

(Continued)
| Study                          | Study design | Number of participants | Follow-up | Outcome measures                                                                 | Results: efficacy for pain intensity                                                                 | Results: SAEs                                                                                     | Results: any other safety data |
|-------------------------------|--------------|------------------------|-----------|----------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|
| Liem et al (2015)             | BA           | Permanent implantation: 32 | 12 months | • AE rate and paresthesia generation  
• Pain intensity by VAS  
• Quality of life by EQ-5D  
• Mood by POMS  
• Physical functioning by BPI | Baseline VAS (N=32): 77.6±2.1  
VAS at 12 months (N=25): 33.6±6.3; P<0.005  
From baseline to 12 months follow-up overall pain improved by 56.3%±8.4%  
The proportion of subjects achieving at least 50% improvement of their overall pain was 60% | 9 SAEs occurred of which 3 (37.5%) related to device  
86 safety events reported across 29 subjects |
| van Bussel et al (2018)       | BA           | 12                     | 16 days trial period (at 3 time points) 1, 3, 6, and 12 months | • Patients’ preference for one stimulation method over another  
• Pain intensity by VAS  
• Condition by GPE scale | Baseline VAS: 68  
Reduction of pain due to DC or DRG stimulation were comparable between the participants | No SAEs occurred  
3 AEs experienced by 2 participants due to device or surgical procedure |
| Goebel et al (2018)           | CR           | 1                      | 1 and 17 months | • Pain intensity by NRS  
• Disability by ODI  
• Interface of pain with daily activities by BPI | Baseline NRS: 7–8  
NRS at 1 month: 25% pain relief  
NRS at 17 months: 60% pain relief with device turned on, and 24-hour pain intensity of 3–4 | Not reported  
Not reported |
| Skaribas et al (2019)         | CR           | 5                      | 1, 2, 3, and 6 months | • Pain intensity by NRS  
• Function by NRS  
• Use of pain medication | Baseline NRS: 8–10  
NRS at trial period: 0–3 | Not reported  
Not reported |
| van Bussel et al (2015)       | CR           | 1                      | 3 months     | • Pain intensity by NRS | Baseline NRS: 6–9  
NRS at 8 days: 1  
NRS at 1 and 3 months: 1–2 | Not reported  
Not reported |

(Continued)
| Study                  | Study design | Number of participants | Follow-up                                      | Outcome measures                                      | Results: efficacy for pain intensity | Results: serious adverse events | Results: any other safety data |
|-----------------------|--------------|------------------------|------------------------------------------------|-------------------------------------------------------|-------------------------------------|---------------------------------|---------------------------------|
| van Buyten et al (2015) | CR           | 11                     | Trial period: 11 and 1 week, 1 month, 5 weeks, 2, 3, 6, and 12 months | • Pain intensity by VAS                                  | Baseline VAS: 77.9±4.2              | No SAEs occurred                  | No other complications occurred |
|                       |              |                        |                                                | • Impact of pain by BPI                                 | VAS at trial period: 14.0±4.2, 81.9% reduction |                                |                                  |
|                       |              |                        |                                                | • Mood by POMS                                         | VAS at 1 week after INS: 27.1±7.6 mm, reduction by 65.2±10.3%, P<0.001 |                                |                                  |
|                       |              |                        |                                                | • Quality of life by EuroQol 5                          | VAS at 1 month: 30.0±10.0, reduction by 62.1%, P<0.005 |                                |                                  |
|                       |              |                        |                                                | • AEs                                                 | VAS at 3 months (n=7): 26.1±11.6, P<0.001, reduction by 63.1%±13.2%, P<0.005 |                                |                                  |
|                       |              |                        |                                                |                                                       | VAS at 6 months: 29.4±11.3, reduction by 63.1%±13.2%, P<0.005 |                                |                                  |
|                       |              |                        |                                                |                                                       | VAS at 12 months: 30.3±12.7, reduction by 61.7%±16.4%, P<0.05 |                                |                                  |
|                       |              |                        |                                                |                                                       | 5 of the 7 subjects (71.4%) had ≥50% pain relief     |                                |                                  |
| Yang and Hunter (2017) | CR           | 2                      | 8 months                                       | • Pain relief by NRS                                    | Patient 1: baseline NRS: 8–9 VAS at 8 months: 90% pain relief | Not reported                     | Not reported                     |
| Low back pain         |              |                        |                                                |                                                       | Patient 2: baseline NRS: 8–9 VAS at 8 months: 5       |                                |                                  |
| Deer et al (2013)     | BA           | 10                     | 3–7 days (minimum of 3 days) postoperatively and last follow-up | • Pain intensity by VAS                                 | Baseline VAS (8 subjects): 73±10                  | 3 SAEs reported, none of which related to device | 14 events reported in 6 subjects; 12 were related to device and included device inactivation, lead migration, and one possible reaction to antibiotic |
|                       |              |                        |                                                | • Perceived percentage of pain relief at the final follow-up visit | VAS postoperatively: 59±17, reduction by 18% |                                |                                  |
|                       |              |                        |                                                | • Improvement in condition by 11-point Likert scale     | VAS at last follow-up: 24±18, reduction by 67%       |                                |                                  |
|                       |              |                        |                                                | • Physicians rated Global Impression of Change on 7-point scale | The average reduction in pain between baseline and final visit was 70 %±22% (P<0.0007) with 88% experiencing at least 30% reduction and 75% reported at least 50% reduction in pain |                                |                                  |
| Study (Continued) |
|-------------------|
| **Table 1** |
| **Study** | **Study design** | **Number of participants** | **Follow-up** | **Outcome measures** | **Results: efficacy for pain intensity** | **Results: serious adverse events** | **Results: any other safety data** |
| Huygen et al (2018)$^d$ | BA | 12 | 1 week, 1, 3, 6, and 12 months | • Pain intensity by VAS  
• Physical functioning by BPI  
• Mood by POMS  
• Quality of life by EQ-SD  
• AEs | Baseline overall VAS: 77.6±2.0  
VAS at 6 months: 49.2±9.6, reduction by 38.2%  
VAS at 12 months: 44.6±9.5, reduction by 44.2%, \( P < 0.001 \)  
Baseline LBP VAS: 73.9±3.7  
VAS at 6 months: 37.5±9.4, reduction by 50.8%  
VAS at 12 months: 40.4±9.7, reduction by 45.5%, \( P < 0.001 \) | Total of 4 SAEs (33.3%): temporary loss of leg strength following the procedure, postdural puncture headache, bladder infection, depression.  
2 AEs (16.7%): discomfort at the implantation site, wound infection plus 4 lead revisions (25.0%) recorded across 12 subjects |
| Huygen et al (2019)$^e$ | BA | 66  
INS: 56 | 1 week, 1, 3, 6, and 12 months | • Pain intensity by VAS  
• Physical functioning by BPI  
• Mood by POMS  
• Quality of life by EQ-SD  
• AEs | Baseline VAS: 8  
VAS at 12 months: 4.1  
24/49 (49%) had \( \geq 50\% \) reduction  
VAS at 12 months:  
FBSS: average reduction in VAS of 54.7±36.9%, \( N=22 \)  
Causalgia: average reduction of 43.7±69.26%, \( N=10 \)  
CRPS: reduction of 46.8±33.9%, \( N=9 \) | Total of 15 SAEs in 14 subjects:  
One death due to medication overdose, implant site infection, INS pocket infection, transient motor deficit, dural puncture, bladder infection, pain following a capsaicin (Qutenza) application, perianal fistula, knee cyst, transient ischemic attack, worsening of preexisting CRPS, bowel obstruction  
Total of 9 non-SAEs in 8 subjects: Loss of stimulation and increased pain, pain at the implant site, implant site wound infection, postimplantation headache, knee pain, depression, accidental burn wound legs |

(Continued)
Table 1 (Continued)

| Study | Study design | Number of participants | Follow-up | Outcome measures | Results: efficacy for pain intensity | Results: serious adverse events | Results: any other safety data |
|-------|--------------|------------------------|-----------|------------------|-------------------------------------|---------------------------------|-------------------------------|
| Weiner et al (2016)\(^{27}\) FBSS low back pain (chronic intractable neuropathic trunk and/or lower limbs pain) | BA | 11 | 2, 4, and 6 weeks | - Pain intensity by VAS  
- Stimulation amplitude  
- AEs | 50% reduction in VAS: in 7/11 participants (63%)  
25%–50% reduction in VAS: in 2/11 participants  
0%–25% reduction in VAS: in 2/11 participants  
The average overall pain reduction was 59.9%, regardless of device placement | No SAEs occurred | No other complication observed |
| Billet et al (2017)\(^{38}\) | CR | 1 | 3 and 5 weeks, and 2, 3, and 6 months | - Pain intensity by VAS  
- ODI  
- Quality of life by EQ-5D-5L  
- PGIC | 66% improvement for back pain and 56% for leg pain at 6-month follow-up | Not reported | Not reported |
| Billet et al (2018)\(^{39}\) Chronic intractable neuropathic pain of the trunk and/or lower limbs due to FBSS | CR | 8 | 2 weeks, 1, 2, and 3 months | - Pain intensity by VAS  
- ODI  
- PGIC  
- Medication usage | Back pain baseline VAS: 71  
Back pain VAS at 3 months: 28  
Leg pain baseline VAS: 28  
Leg pain VAS at 3 months: 13 | One stimulator migration that required revision | Not reported |
| Morgalla et al (2018)\(^{40}\) Groin pain | BA | 34 | At 3 years follow-up: 11 | - Pain intensity by VAS  
- PDI  
- PCS  
- BPI  
- BDI | Baseline VAS: Mdn =8  
VAS at 3 months: Mdn =3  
VAS at 1 year: Mdn =3.5  
VAS at 2 years: Mdn =4  
VAS at 3 years: Mdn =4.5  
Significant decrease in pain after 3 months, 1 and 2 years (P<0.001), and after 3 years (P<0.005) when compared to the baseline measurement | Five subjects showed complications (16.7%)  
Breakage of the lead, an infection during the test trial, generator needed relocation, an additional electrode required | Not reported |
| Study                          | Study design | Number of participants | Follow-up                              | Outcome measures                          | Results: efficacy for pain intensity                  | Results: serious adverse events | Results: any other safety data |
|-------------------------------|--------------|------------------------|----------------------------------------|-------------------------------------------|-------------------------------------------------|---------------------------------|--------------------------------|
| Schu et al (2015)§            | CS           | 25                     | Average follow-up period was 27.8±4.3 (SEM) weeks; median: 26.0 weeks, range: 0–68 weeks | • Pain intensity by VAS                   | Baseline VAS: 74.5±1.8 VAS at follow-up (n=23): 20.7±3.9, a mean improvement of 71.4±5.6% >80% reduction in VAS; in 47.8%; or 11/23 of participants >50% reduction in VAS; in 82.6% or 19/23 of participants | Not reported                     | Not reported                    |
| Zuidema et al (2014)§         | CR           | 3                      | 1 week, 1, 3, 6, and 12 months (data given in results are just for the follow-up period of 2 and 3 months) | • Pain intensity by VAS • Quality of life | Patient 1: baseline VAS: 90 VAS at 2 weeks and 3 months: 0 Patient 2: baseline VAS: 90 VAS at 1 week and 2 months: 10 Patient 3: baseline VAS: 95 VAS at 1 week and 2 months: 10 No SAEs occurred | No other complications observed |                                |
| Pelvic girdle pain            |              |                        |                                        |                                            |                                                  |                                 |                                |
| Hunter and Yang (2019)§       | CR           | 7                      | 3–12 months, different for each patient | • Pain intensity by NRS                  | Reported in figures of a manuscript Patient 1: baseline NRS: 7–9 NRS at ~12 months: 2 Patient 2: baseline NRS: 8–9 NRS at 7 months: 0 Patient 3: baseline NRS: 4–8 NRS at 7 days trial: 1–2 Patient 4: baseline NRS: 7–8 NRS at 3 days trial: 60% reduction in pain Patient 5: baseline NRS: 6 NRS at 12 months: 2 Patient 6: baseline NRS: 6, pain was reduced with different treatments, but returned after some time (NRS unknown) NRS at 7 days trial: 0 Patient 7: baseline NRS: 9 At 7 days trial 85% reduction in pain | Not reported                     | Not reported                    |

(Continued)
Table 1 (Continued)

| Study | Study design | Number of participants | Follow-up | Outcome measures | Results: efficacy for pain intensity | Results: serious adverse events | Results: any other safety data |
|-------|--------------|------------------------|-----------|-----------------|-------------------------------------|---------------------------------|-------------------------------|
| Rowland et al (2016)\(^a\) | CR           | 1                      | 6 months  | • Pain intensity by NRS  
• MPQ     | Baseline NRS: 7 and 10 at worst  
NRS at 6 months: 4 (43% pain reduction) | Not reported | Not reported |
| Peripheral neuropathy including diabetic peripheral neuropathy | | | | | | | |
| Eldabe et al (2018)\(^a\) | CS           | 10                     | At 12 months follow-up: 5  
1 week, 1, 3, 6, and 12 months | • Pain intensity by VAS  
• Device complications and procedure-related AEs | Baseline VAS: 79.6±13.5  
VAS after permanent implantation: 48.6±16.1, reduction by 63.90±21.4%  
VAS at 6 months: 49.4±9.2, reduction by 65.6±27.1%  
VAS at 12 months: 48.3±25.3 reduction by 64.2±35.8% | Lead dislodgement required system to be explanted | Postdural puncture headache related to the procedure |
| Maino et al (2017)\(^a\) | CR           | 1                      | 10 days, 2, 6, 12, and 20 months | • Pain intensity by NRS  
• MPQ  
• Degree of disability by ODI | Baseline NRS: 8 at rest and 2–3 during the day  
NRS during 10 days of trial period: 3 at rest (improvement of 62.5%). NRS at 2 months: 2 (improvement of 75%)  
NRS at the 6 months: 2  
NRS at 12 and 20 months: 4 | Not reported | Not reported |
| van Velsen et al (2018)\(^a\)  
Intractable foot pain – idiopathic peripheral neuropathy | CR           | 1                      | 7 days and 2 months | • Pain intensity by VAS  
• Function  
• Use of pain medication | Baseline VAS: 9  
VAS at 7 days: 1, reduction by 85%  
VAS at 2 months: 1 | No SAEs occurred | No other complications were reported |
| van Velsen et al (2018)\(^a\) | CR           | 1                      | 2 months  | • Pain intensity by VAS | Baseline VAS: 9;  
VAS after 7-day trial: 1;  
VAS at 2 months: 1, reduction by 85% | No SAEs occurred | No other complications observed |

(Continued)
Table 1 (Continued)

| Study | Study design | Number of participants | Follow-up | Outcome measures | Results: efficacy for pain intensity | Results: serious adverse events | Results: any other safety data |
|-------|--------------|------------------------|-----------|------------------|--------------------------------------|---------------------------------|------------------------------|
| Phantom limb pain | | | | | | | |
| Eldabe et al (2015) | CR | 8 | 9±6.3 months (from 5 to 24 months) | • Pain intensity by VAS  
• Quality of life by EQ-SD  
• Medication use  
• Success was defined as 50% or greater pain relief | Baseline VAS: 83.5±10.5  
VAS at last follow-up: 38.9±27.1  
The percentage of pain reduction was on average 52±3.9  
In 3 of 8 participants pain relief diminished over time | No SAEs occurred | No other complications were reported |
| Chronic intractable pain in the coccyx | | | | | | | |
| Giordano et al (2018) | CR | 1 | 4 months | • Pain intensity by VAS | Baseline VAS: 8  
VAS at 4 months: 90% pain reduction | No SAEs occurred | No other complications were reported |
| Chronic testicular pain | | | | | | | |
| Hassanain and Murphy (2019) | CR | 1 | 1 year | • Pain intensity by VAS | Baseline VAS: not reported  
VAS at 1 year: sustained pain relief of 70%-80% | Not reported | Not reported |
| Anterior cutaneous nerve entrapment syndrome | | | | | | | |
| Mol and Roumen (2018) | CR | 5 | 6 and 12 months | • Pain intensity by NRS  
• Medication use | Patient 1: baseline NRS: 8  
NRS at 12 months: 0  
Patient 2: baseline NRS: 9  
NRS at 1 month: 5  
NRS at 6 months: 4  
Patient 3: baseline NRS: 9  
NRS at 6 months: 9  
Patient 4: baseline NRS: 8  
NRS at 6 months: 6  
NRS at 12 months: 3  
Patient 5: baseline NRS: 8  
NRS at 6 months: 8 | Lead dislocation, lead breakage, pain at the battery site, and overstimulation in 4 subjects | Not reported |
| Loin pain hematuria syndrome | | | | | | | |
| Zuidema and Schapendonk (2017) | CR | 1 | 2 weeks, 1 month, 5 weeks, 6 months, and 3 years | • Pain intensity by NRS  
• Use of pain medication | Baseline NRS: 9  
NRS at 1 and 6 months: 1  
NRS at 3 years: 3–4 | Lead migration and stimulator movement | No other complications observed |

Abbreviations: AEs, adverse events; BA, before and after comparison; BDI, beck depression inventory; BPI, brief pain inventory; CBA, controlled before and after; CR, case report; CRPS, complex regional pain syndrome; CS, case series; DC, dorsal column; DRG, dorsal root ganglion; EQ-SD, EuroQol five-dimensions questionnaire; FBSS, failed back surgery syndrome; GPE, global perceived effect; INS, permanent DRG stimulator implanted; LBP, low back pain; MPQ, McGill Pain Questionnaire; NRS, Numeric Rating Scale; ODI, Oswestry Disability Index; PDI, Pain Disability Index; PCS, pain catastrophizing scale; POMS, Profile Of Mood States; PRS, pain relief scale; RCT, randomized controlled trial; SAEs, serious adverse events; SCS, spinal cord stimulation; SEM, standard error of mean; SF-36, short form health survey; TNS, trial period of DRG stimulator implantation; VAS, visual analog scale.
Table 2 Inclusion and exclusion criteria and baseline characteristics of participants

| Study                      | Inclusion criteria/previous treatment                                                                 | Exclusion criteria                                                                                      | Baseline characteristics                                      |
|----------------------------|-------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------|---------------------------------------------------------------|
| Deer et al (2017)          | • CRPS and/or peripheral causalgia for at least 6 months with chronic, intractable pain              | • Back pain was the greatest region of pain                                                             | DRG arm:                                                     |
|                            | • Age between 22 and 75 years                                                                       | • Pregnant or nursing, plans to become pregnant                                                       | • 51.3% of females                                           |
|                            | • Naive to stimulation                                                                              | • Escalating or changing pain condition 30 days prior to study enrollment                              | • 94.7% white race                                          |
|                            | • Minimum baseline VAS 60 mm in the area of greatest pain                                             | • Involved in medically related litigation                                                            | • Average age 52.4 years                                     |
|                            | • Failed at least 2 prior pharmacologic treatments from 2 different drug classes                     | • Corticosteroid therapy at an intended site of stimulation 30 days or RF 3 months prior to study enrollment | • Average body mass index 30.5 kg/m²                          |
|                            | • Stable neurologic function 30 days prior to screening                                              | • Pain medication(s) dosage(s) was not stable for at least 30 days prior to study enrollment           | • Average duration of chronic lower limb pain 7.5 years      |
|                            | • Free from psychological pathology that contraindicated an implantable device                       | • Previously failed SCS therapy                                                                       | SCS arm:                                                     |
|                            |                                                                                                      | • An active implantable device                                                                       | • 51.3% of females                                           |
|                            |                                                                                                      | • Pain only within a cervical distribution                                                            | • 92.1% white race                                          |
|                            |                                                                                                      | • Cognitive, physical, or sensory impairment                                                           | • Average age 52.5 years                                     |
|                            |                                                                                                      | • An indwelling device                                                                                 | • Average body mass index 28.9 kg/m²                          |
|                            |                                                                                                      | • An active systemic infection                                                                       | • Average duration of chronic lower limb pain 6.8 years      |
|                            |                                                                                                      | • Medical comorbidity that contraindicates placement of device                                         | Comorbidities and medications taken for subject conditions were similar in both arms. No statistically significant differences were found among the baseline characteristics between treatment arms |
|                            |                                                                                                      | • Participation in another clinical investigation with 30 days prior to study enrollment               |                                                               |
|                            |                                                                                                      | • Coagulation disorder or uses anticoagulants                                                        |                                                               |
|                            |                                                                                                      | • Diagnosed with cancer within 2 years prior to inclusion                                             |                                                               |
|                            |                                                                                                      | • Imaging findings within 12 months prior to study enrollment                                         |                                                               |
|                            |                                                                                                      | • Presence of an escalating or changing pain condition within the month prior to enrollment           |                                                               |
|                            |                                                                                                      | • Pain only within a cervical distribution                                                            |                                                               |
|                            |                                                                                                      | • Corticosteroid therapy at an intended site of stimulation within the 30 days or RF treatment within the 3 months prior to study enrollment |                                                               |
|                            |                                                                                                      | • Had a coagulation disorder                                                                         |                                                               |
|                            |                                                                                                      | • Had an indwelling device                                                                             |                                                               |
|                            |                                                                                                      | • Had an active implantable device                                                                     |                                                               |
| Liem et al (2013)          | • Chronic, intractable pain in the trunk, limbs, and/or sacral region for at least 6 months         |                                                                                                        | 17 females and 15 males                                     |
|                            | ±18 years old                                                                                        |                                                                                                        | Mean age of men 58.9±8.9 years                              |
|                            | • Minimum baseline VAS 60 mm                                                                          |                                                                                                        | Mean age of women 46.9±12.5 years                            |
|                            | • Failed other treatment modalities (pharmacological and/or surgical)                               |                                                                                                        | Subjects had chronic pain of neuropathic origin of varying etiologies |
|                            | • Have stable pain medication dosage for a minimum of 30 days prior to study enrollment              |                                                                                                        |                                                               |
|                            | • Have a stable pattern of neurological symptoms                                                      |                                                                                                        |                                                               |
| Liem et al (2015)          | Same as Liem et al (2013)                                                                            | Same as Liem et al (2013)                                                                             | Same as Liem et al (2013)                                   |
| Goebel et al (2018)        | • Ineffective treatment with pamidronate, steroids, opioids                                          |                                                                                                        |                                                               |
|                            | • Failed SCS                                                                                         |                                                                                                        |                                                               |
|                            | • Repeated intensive rehabilitation program with limited success                                      |                                                                                                        |                                                               |
|                            | • Intravenous immunoglobulin over a 6 month period                                                   |                                                                                                        |                                                               |
|                            |                                                                                                      | NA                                                                                                      | Male (age not written)                                      |
| Skaribas et al (2019)      | • Previous back surgery                                                                              |                                                                                                        | CRPS in a period of 6 years prior to DRG stimulation         |
|                            | • Allodynia, hyperpathia, edema, purplish discoloration indicating vasomotor changes, and decreased range of motion of the affected foot |                                                                                                        |                                                               |
|                            |                                                                                                      | NA                                                                                                      | 4 females, 1 male                                          |
|                            |                                                                                                      |                                                                                                        | Age between 49 and 71 years                                 |

(Continued)
| Study                          | Inclusion criteria/previous treatment                                                                 | Exclusion criteria                                                                 | Baseline characteristics       |
|-------------------------------|---------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------|--------------------------------|
| van Buyten et al (2015)35     | Same as Liem et al (2013)3                                                                           | Same as Liem et al (2013)3                                                          | 6 females, 2 males             |
|                               | • 6 females, 2 males                                                                                   | • Average age 43.9±5.6 years                                                        | Average age 43.9±5.6 years     |
|                               | • Average age 43.9±5.6 years                                                                           | • Range: 18–65 years                                                                | Range: 18–65 years             |
|                               | • Range: 18–65 years                                                                                  | • VAS score at baseline 77.9±4.2 mm                                                 | VAS score at baseline 77.9±4.2 mm |
| van Bussel et al (2018)31     | • CRPS for a minimum of 1 year or longer                                                               | • Previous neurostimulation                                                         | 11 females, 1 male             |
|                               | • ≥18 years old                                                                                       | • Depression or anxiety disorder                                                    | Mean age 38.7 years (range 22–57 years) |
|                               | • Minimum baseline VAS 50 mm or higher                                                                   | • Pregnancy or pregnancy desire within 1 year                                       | VAS score at baseline 68 mm     |
|                               | • No improvement in symptoms after ≥1 year of treatment according to the Dutch guidelines for CRPS     | • Body mass index >35 kg/m²                                                          | None of the included subjects had demonstrable nerve injury in the affected knee |
|                               |                                                                                                        | • Life expectancy                                                                    |                                |
|                               |                                                                                                        | • Participants unable to complete the questionnaires                                 |                                |
|                               |                                                                                                        | • An active implantable device                                                       |                                |
|                               |                                                                                                        | • Anticoagulant drug therapy or disturbed coagulation                               |                                |
|                               |                                                                                                        | • Immunocompromised participants                                                    |                                |
|                               |                                                                                                        | • Drugs/medication/alcohol addiction                                                |                                |
|                               |                                                                                                        |                                                                                     |                                |
|                               | • 11 females, 1 male                                                                                  |                                                                                     |                                |
|                               | • Mean age 38.7 years (range 22–57 years)                                                              |                                                                                     |                                |
|                               | • VAS score at baseline 68 mm                                                                          |                                                                                     |                                |
|                               | • None of the included subjects had demonstrable nerve injury in the affected knee                       |                                                                                     |                                |
| van Bussel et al (2015)32     | • Extensively treated with different types of oral medication                                          | NA                                                                                  | 48 years old women             |
|                               | • Lumbar sympathetic block resulted in no clinically significant relief of symptoms                     |                                                                                     | 5 years of symptoms            |
|                               | • Physical therapy failed                                                                             |                                                                                     |                                |
| Yang and Hunter (2017)33      | • Failed SCS                                                                                            | NA                                                                                  | 43-year-old female             |
|                               |                                                                                                        |                                                                                     | 2 years of symptoms            |
|                               |                                                                                                        |                                                                                     | Baseline NRS 8–9               |
|                               |                                                                                                        |                                                                                     | 50-year-old female             |
|                               |                                                                                                        |                                                                                     | 9 years of symptoms            |
|                               |                                                                                                        |                                                                                     | Baseline NRS 8–9               |
| Low back pain                 |                                                                                                        |                                                                                     |                                |
| Deer et al (2013)38           | • Chronic, intractable neuropathic pain of the trunk and/or limbs                                       | • Previous posterior fusion                                                          | 5 females, 5 males             |
|                               | • ≥18 years old                                                                                       | • Severe foraminal stenosis at the expected target level                             | Average age of men 52±5 years   |
|                               | • Failed other treatment modalities including opioid and nonopioid-based pain medication, physical therapy, epidural steroid injections, selective nerve root blocks, trigger point injections, medial branch radiofrequency, intrathecal pump implantation, and SCS | • Presence of indwelling implantable devices such as cardiac devices, spinal cord or peripheral nerve stimulators, or vascular access catheters | Average age of women 39±4 years |
|                               | • No change in medications, surgery, injections, or other treatment for a minimum of 30 days prior to study enrollment | • Pregnancy                                                                         |                                |
|                               | • Consistent pattern of pain and neurological symptoms for a minimum of 30 days prior to study enrollment | • Pain patterns could not be >50% in the axial spine, but axial spine participants were not excluded |                                |

(Continued)
Table 2 (Continued)

| Study                  | Inclusion criteria/previous treatment                                                                                                                                                                                                 | Exclusion criteria                                                                                                                                                                                                 | Baseline characteristics                                                                                     |
|-----------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------|
| Huygen et al (2018)   | • Back pain due to FBSS  
• Axial LBP as either primary or secondary region of pain  
• Minimum baseline VAS ≥60 mm  
• Successful DRG stimulation trial with better than 50% pain relief  
• At least one lead permanently implanted at an L2 or L3 DRG | Not reported                                                                                                                                                                                                           | • 12 participants, 33% male  
• Average age 51.1 years  
• Baseline VAS 77.6 ±2.0 mm  
• All subjects had also leg pain  
• 6 subjects had foot pain  
• 2 subjects had buttock pain |
| Huygen et al (2019)   | • ≥18 years old  
• Chronic pain for at least 6 months  
• Minimum baseline VAS of ≥60  
• Pain limited to the lower body: in the thoracic, lumbar, and/or sacral distributions  
• Failed other treatment modalities including pharmacological therapy, physical therapy, and interventional pain procedures for chronic pain  
• Psychologically appropriate for the implantation | • Pain primarily in cervical distribution  
• Unstable pain condition  
• Corticosteroid or radiofrequency treatment at the intended site of stimulation prior to study enrollment  
• Presence of an active implantable device  
• Coagulation disorder or use of anticoagulants  
• Cancer  
• Pregnancy | • 64% females  
• Average age 52 ±1.5 years  
• Range from 30 to 80 years |
| Weiner et al (2016)   | • Chronic intractable neuropathic pain of the trunk and/or lower limbs due to FBSS  
• ≥18 years old  
• Minimum baseline VAS of >5  
• Speaking English or Spanish  
• Pass a psychological evaluation  
• Have the cognitive ability to use the external transmitter | • Visceral pain  
• Hyperalgesia or allodynia of the lower back  
• Allergies to system components  
• Active cancer treatment  
• Drug dependence  
• Pregnancy  
• Inability to comply with the study requirements | • 55% female  
• Mean age 63 years |
| Billet et al (2017)   | • Microdisectomy with no results  
• Anterior lumbar interbody fusion performed  
• Medical management with tramadol HCl and paracetamol  
• PRF treatment followed by ablation of the facet joint did not provide pain relief | NA                                                                                                                                                                                                                 | • Patient had traffic accident resulting in traumatic disc herniation |
Table 2 (Continued)

| Study                          | Inclusion criteria/previous treatment                                                                 | Exclusion criteria                                                                                      | Baseline characteristics                                      |
|-------------------------------|-------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------|---------------------------------------------------------------|
| Billet et al (2018)39         | • Chronic intractable neuropathic pain of the trunk and/or lower limbs due to FBSS                   | • Malignancies, postherpetic neuralgia, active systemic infection                                       | • 2 females, 4 males                                         |
|                               | • ≥18 years old                                                                                       | • Immune-compromised, insulin dependent                                                                  | • Mean age 53 years                                           |
|                               | • Minimum baseline VAS >50 mm                                                                          | • Diabetes not controlled through diet and/or medication                                                 | • Range 33–67 years                                           |
|                               | • Refractory to conventional medical management                                                        | • Bleeding complications, coagulopathy issues                                                           | • Four subjects reported also leg pain                       |
|                               | • Speaking Dutch or French                                                                            | • Life expectancy of <1 year                                                                            |                                                               |
|                               | • Pass a psychological evaluation                                                                     | • Active implanted device                                                                               |                                                               |
|                               | • Have the cognitive ability to use the external transmitter                                            | • Pregnancy                                                                                                |                                                               |
|                               | • Live within a radius of 75 km                                                                       | • Inability to comply with the study requirements                                                       |                                                               |
|                               | • 2 females, 4 males                                                                                   |                                                                                                        |                                                               |
|                               | • Mean age 53 years                                                                                     |                                                                                                        |                                                               |
|                               | • Range 33–67 years                                                                                    |                                                                                                        |                                                               |
|                               | • Four subjects reported also leg pain                                                                   |                                                                                                        |                                                               |
|                               | • 2 females, 4 males                                                                                   |                                                                                                        |                                                               |
|                               | • Mean age 53 years                                                                                    |                                                                                                        |                                                               |
|                               | • Range 33–67 years                                                                                    |                                                                                                        |                                                               |
| Groin pain                    |                                                                                                       |                                                                                                        |                                                               |
| Morgalla et al (2018)41       | • Chronic neuropathic pain in the groin                                                                | • Previous spinal surgery at the level of the intended implantation of the DRG leads                   | • 13 females, 21 males                                       |
|                               | • Pain confirmed by a clinically detectable sensory loss, hyperalgesia, or allodynia, within an anatomic   | • Cardiac pacemakers, vascular access catheters, other spinal cord stimulators, or peripheral nerve      | • Mean age 50.4 years                                        |
|                               | concordant area of a nerve or a root dermatome                                                          | stimulators (PNSs)                                                                                      | • Range 24–84 years                                          |
|                               | • Failure of pain treatment using various medication, interventions, or even hospitalization              | • Psychiatric disorders including anxiety and depression                                               | • History of pain for longer than 6 months                   |
|                               | • No further indication for another surgical intervention in the area of the previously operated groin   |                                                                                                        | • Mean duration of pain 2.5 years, from 0.5 to 8 years        |
| Schu et al (2015)8            | • Chronic, intractable neuropathic pain of the groin                                                    | • Previous posterior fusion                                                                            |                                                               |
|                               | • ≥18 years old                                                                                       | • Severe foraminal stenosis at the expected target level                                                |                                                               |
|                               | • Failed other treatment modalities including oral medications and/or interventional procedures or    | • Presence of current indwelling implantable devices                                                    |                                                               |
|                               | surgical intervention                                                                                  | • Pregnancy                                                                                                |                                                               |
|                               | • Mean baseline VAS (N=25) 74.5±1.8 mm                                                                    |                                                                                                        |                                                               |
|                               | • The most frequent diagnosis was herniorrhaphy (N=12)                                                  |                                                                                                        |                                                               |
|                               | • Other subjects had a variety of pain etiologies, many related to postsurgical pain                   |                                                                                                        |                                                               |
|                               | • No data about age or sex                                                                             |                                                                                                        |                                                               |
| Zuidema et al (2014)7         | • Pain refractory to antineuropathic medication (pregabalin and amitriptyline)                         |                                                                                                        |                                                               |
|                               | • No results with TENS and PRF and local corticoid infiltration                                         |                                                                                                        |                                                               |
|                               |                                                                                                       |                                                                                                        | • 36 years old men                                           |
|                               |                                                                                                       |                                                                                                        | • 5 years of chronic pain, baseline VAS 90 mm               |
|                               |                                                                                                       |                                                                                                        | • 39 years old female, 6 years of chronic pain, baseline   |
|                               |                                                                                                       |                                                                                                        | • VAS 90 mm                                                  |
|                               |                                                                                                       |                                                                                                        | • 46 years old female, 4 years of chronic pain, baseline   |
|                               |                                                                                                       |                                                                                                        | • VAS 95 mm                                                  |
| Pelvic girdle pain            |                                                                                                       |                                                                                                        |                                                               |
| Hunter and Yang (2019)9       | • Pain refractory to medication, neurolysis, surgery, and, in some cases, SCS                          |                                                                                                        | • 4 females, 3 males                                         |
|                               | • Variation in location and presentation of pain, suspected cause/etiology, and associated symptoms    |                                                                                                        | • Age range from 36 to 63 years                              |
|                               |                                                                                                       |                                                                                                        |                                                               |
|                               |                                                                                                       |                                                                                                        | (Continued)                                                  |
Table 2 (Continued)

| Study                        | Inclusion criteria/previous treatment                                                                 | Exclusion criteria                                                                 | Baseline characteristics                  |
|------------------------------|--------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------|-------------------------------------------|
| Rowland et al (2016)         | • Failed trials of physiotherapy, gabapentin, and steroid injections                                 | NA                                                                                  | • 37-year-old female                     |
|                              | • On the day of admission, the patient received oral treatment with paracetamol 1 g four times daily |                                                                                     | • 9 years of chronic pain                |
|                              | • Diazepam 5 mg, MST 60 mg twice daily, Zomorph (morphine sulfate) 10 mg twice daily and amitriptyline 25 mg once daily |                                                                                     | • Baseline NRS 7                         |

Peripheral neuropathy including diabetic peripheral neuropathy

| Study                        | Inclusion criteria/previous treatment                                                                 | Exclusion criteria                                                                 | Baseline characteristics                  |
|------------------------------|--------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------|-------------------------------------------|
| Eldabe et al (2018)          | • Chronic intractable pain due to diabetic polyneuropathy of the lower limbs for at least 6 months     | • Unstable neurological symptoms                                                    | • 10 male diabetic participants           |
|                              | • ≥18 years old                                                                                       | • A baseline VAS score of <60 mm                                                    | • Mean age 65.2 [SD 8.8] years            |
|                              | • Stable pain medication for minimum 30 days prior to study enrollment                                | • Pregnancy                                                                         | • Mean duration of PDPN-related symptoms 7.0 (SD 3.1) years, ranging from 3 to 11 years (n=5) |
|                              | • Failed previous interventions including SCS                                                       | • Implanted neuromodulation devices                                                 | • Average baseline VAS 79.6 (SD 13.5) mm   |
|                              | • Patient primary pain area was considered any part of the lower limbs including leg, thigh, shin, calf, and foot |                                                                                     |                                           |
| Maino et al (2017)           | • No results with multiple medications including gabapentin, pregabalin, duloxetine, amitriptyline, mirtazapine, lidocaine patches, topical capsaicin 8%, and cannabis | NA                                                                                  | • 74-year-old men                        |
|                              | • TENS, physical therapy, acupuncture, and a corticosteroid infiltration with no results             |                                                                                     | • Baseline pain NRS 8                    |
|                              | • Ice packs and pressure applied on left foot provided some relief at night                           |                                                                                     | • Chronic pain for 6 years               |
|                              |                                                                                                       |                                                                                     | • Hyperlipidemia, left common carotid artery stenosis, coronary artery disease, and a depressive disorder |
| van Velsen et al (2018)      | • Use of neuropathic pain medications such as gabapentin, pregabalin, topiramate, and duloxetine and opioid analgesics such as hydrocodone, tapentadol, oxycodone, and methadone with no results | NA                                                                                  | • Oral daily doses of aspirin 100 mg and atorvastatin 20 mg |
|                              | • Traditional SCS did not give satisfactory pain relief                                              |                                                                                     |                                           |
| van Velsen et al (2018)      | Same as van Velsen et al (2018)                                                                       | NA                                                                                  | • 45-year-old Caucasian male patient      |
|                              |                                                                                                       |                                                                                     | • Chronic pain for 2 years               |

(Continued)
### Table 2 (Continued)

| Study                          | Inclusion criteria/previous treatment | Exclusion criteria | Baseline characteristics |
|-------------------------------|---------------------------------------|-------------------|--------------------------|
| **Phantom limb pain**         |                                       |                   |                          |
| Eldabe et al (2015)44         | Not reported                          | Not reported      | • 5 females and 3 males  |
|                               |                                       |                   | • Age range from 28 to 76 years |
|                               |                                       |                   | • Baseline medication use including clonidine, bupivacaine, morphine, pregabalin, oramorph, amitriptyline, tramadol, zomorph, Lansoprazol, targin, amineurin, ariclaim, lyrica, palesia, clonazepam, oxycodone, gabapentin, trypotizol, and fentanyl patch |
| Giordano et al (2018)45       | • Multiple coccygeal blocks, trigger point injections, epidural steroid injections with no results | NA                | • 37-year-old-female |
|                               |                                       |                   | • 8 years of chronic pain |
|                               |                                       |                   | • Medication regimen consisting of oxycodone 10 mg PO BiD, dexketoprofen 25 mg PO QID, duloxetine 60 mg PO QD, trazodone 100 mg PO QD, and pregabalin 75 mg PO BiD |
| Hassanain and Murphy (2019)46 | • Trial of antineuropathic medications in the form of a combination of amitriptyline and gabapentin | NA                | • 45-year-old men |
|                               | • PRF trial for 3 months (provide 60%–70% pain relief) |                   | • History of obesity, type II diabetes mellitus and obstructive sleep apnea |
| Mol and Roumen (2018)47       | • Patient 1: failed neurectomy, resection of neuroma, use of opiates | NA                | • 35-year-old female, BMI 35.4, chronic pain for 6 years, baseline pain NRS 8 |
|                               | • Patient 2: failed neurectomy, use of pain medications |                   | • 26-year-old female, BMI 26.7, chronic pain for 6 years, baseline pain NRS 9 |
|                               | • Patient 3: use of pain medication |                   | • 50-year-old men, BMI 25.7, duration of pain not given, baseline pain NRS 9 |
|                               | • Patient 4: infiltration with lidocaine, neurectomy |                   | • 18-year-old female, BMI 20.5, baseline NRS 8 |
|                               | • Patient 5: neurectomy and TENS |                   | • 60-year-old men, BMI not given, chronic pain for 2 years, baseline NRS 8 |
| Zuidema and Schapendonk (2017)48 | • Use of antineuropathic analgesics (pregabalin, amitriptylin, and duloxetine) | NA                | • 37-year-old women |
|                               | • Use of opioid and nonopioid analgesics (paracetamol, oxycodone, and tramadol) |                   | • Chronic pain for 2 years |
|                               | • Use of perindopril |                   | • Patient history: urolithiasis and tonsillectomy |
|                               | • Minimal invasive techniques (quadratus lumborum block, repeated neurolytic celiac plexus blocks, and splanchnic nerve blocks) with no results |                   | • Allergy to NSAIDs |
|                               | • TENS and catheter-based renal denervation with no results |                   |                          |

**Abbreviations:** CRPS, complex regional pain syndrome; DRG, dorsal root ganglion; FBSS, failed back surgery syndrome; LBP, low back pain; NA, not applicable; NRS, Numeric Rating Scale; NSAID, nonsteroidal anti-inflammatory drug; PRF, pulsed radiofrequency; SCS, spinal cord stimulation; TENS, transcutaneous electrical nerve stimulation; VAS, visual analog scale.
Table 3  Parameters of electrical field stimulation of dorsal root ganglion

| Study                          | Comparator | Parameters of stimulation | Stimulator used (electrode and device)                                                                 | Position of the leads                                                                 |
|--------------------------------|------------|---------------------------|--------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|
| Deer et al (2017)              | SCS        | Pulse width: 3 months, 306.4±148.1 µs (range 30–1,000 µs); 12 months, 289.8±133.8 µs (range 90–1,000 µs) Frequency: 3 months, 20.8±7.1 Hz (range 10–48 Hz); 12 months, 19.0±5.1 Hz (range 10–36 Hz) Amplitude: 3 months, 915.4±822.0 µA (range 75–6,000 µA); 12 months, 827.4±657.1 µA (range 75–4,000 µA) | For DRG stimulation: Axium Neurostimulator System (Spinal Modulation, Inc.) – up to 4 leads implanted; for SCS: commercially available system (Restore Ultra and Restore Sensor; Medtronic, Minneapolis, MN, USA) – up to 2 leads implanted | From T10 to S2 depending on the dermatomal target corresponding to the subjects’ primary region of pain |
| Liem et al (2013)              | No comparator | Pulse width: 362 ms Frequency: 46 Hz Amplitude: 907 µA | Axium neurostimulator and 4 quadrupolar percutaneous leads and wireless programmer devices (Spinal Modulation, Inc.) | According to individual location and distribution of pain (not specified) |
| Liem et al (2015)              | No comparator | Pulse width: 362 ms Frequency: 46 Hz Amplitude: 907 µA | Axium neurostimulator and 4 quadrupolar percutaneous leads and wireless programmer devices (Spinal Modulation, Inc.) | According to individual location and distribution of pain (not specified) |
| van Bussel et al (2018)        | SCS        | Not written in manuscript | Two 4-contact leads and DRG stimulator (St. Jude Medical Inc., Little Canada, MN, USA) | L3 and L4 DRG |
| Goebel et al (2018)            | No comparator | Not written in manuscript | Not written in manuscript | L4 DRG |
| Skaribas et al (2019)          | No comparator | Not written in manuscript | Two quadrupolar DRG electrodes | S1 |
| van Bussel et al (2015)        | No comparator | Pulse width: 170 µs for lead 1 and 2 (L2 and L3) and 160 µs for lead 3 (L42) Frequency: 20 Hz Amplitude: L2, 700 µA; L3, 1,030 µA; L4, 500 µA | Three quadrupolar DRG stimulation leads (refer to Liem et al (2013); without giving company or stimulator details) | L2–L4 |
| van Buyten et al (2015)         | No comparator | Not written in manuscript | Quadrupolar percutaneous leads and Axium stimulator (Spinal Modulation, Inc.) | According to individual location and distribution of pain |
| Yang and Hunter (2017)         | SCS        | Not written in manuscript | Axium stimulator (Spinal Modulation, Inc.) | L3 and L4 |

(Continued)
Table 3 (Continued)

| Study                  | Comparator | Parameters of stimulation | Stimulator used (electrode and device)                                                                 | Position of the leads                                      |
|------------------------|------------|---------------------------|---------------------------------------------------------------------------------------------------------|-----------------------------------------------------------|
| Low back pain          |            |                           |                                                                                                        |                                                            |
| Deer et al (2013)38    | No comparator | Pulse width: 200 µs; Frequency: 68 Hz; Amplitude: 800 µA | Quadrupolar DRG stimulation leads (Spinal Modulation, Inc.) and external stimulator (Spinal Modulation, Inc) | Thoracic, lumbar, and sacral spinal levels                 |
| Huygen et al (2018)60  | No comparator | Pulse width: 269±17.0 µs, range 80–440 µs, Frequency: 21.3±0.6 Hz, range 20–30 Hz; Amplitude: 591.9±50.3 µA, range 1,750–1,130 µA | Axium neurostimulator (Spinal Modulation, Inc.) – up to 4 leads were implanted per subject, leads in bipolar configuration | L2 or L3 DRG. One lead was placed at each of the L1, L4, L5, and S1 DRGs in subjects with foot and buttck pain |
| Huygen et al (2019)40  | No comparator | Not written in manuscript | Axium neurostimulator (Abbott Laboratories)                                                             | Not written in manuscript                                 |
| Weiner et al (2016)37  | No comparator | Pulse width: 500 µs, Frequency: 100 Hz | The Stimwave Freedom SCS System, including stimulator and electrode (Stimwave Technologies Incorporated, Fort Lauderdale, FL, USA) | L1–L5                                                     |
| Billet et al (2017)36  | No comparator | Pulse width: 30 µs, Frequency: 10 kHz, Amplitude: 1.5 and 2.5 mA | Two Freedom 4A electrodes with four contacts and Freedom SCS external device (Stim Relieve LLC)          | L2                                                        |
| Billet et al (2018)39  | No comparator | Pulse width: 10–1,000 µs, Frequency: 2–10,000 Hz, Amplitude: 1–24 mA | Two Freedom 4A electrodes with four contacts; each electrode array contains four contacts (3 mm in diameter with 4 mm spacing) (Stimwave) | T9 and L2                                                 |
| Groin pain             |            |                           |                                                                                                        |                                                            |
| Morgalla et al (2018)40| No comparator | Not written in manuscript | Not written in manuscript                                                                                  | T12, L1, and L2 (mostly combination of L1 and L2)          |
| Schu et al (2015)41    | No comparator | Pulse width: 137 ms, Frequency: 60 Hz, Amplitude: 6.32 mA | Axium neurostimulator system and quadrupolar DRG stimulation leads (Spinal Modulation, Inc.) | T11 up to L3; Subjects received 1, 2, or 3 leads to cover their pain area. All leads were placed unilaterally |
| Zuidema et al (2014)59 | No comparator | Not written in manuscript | Axium stimulator and DRG lead (Spinal Modulation, Inc.)                                                 | T11, T12, and L2                                          |
| Pelvic girdle pain     |            |                           |                                                                                                        |                                                            |
| Hunter and Yang (2019)9 | No comparator | Not written in manuscript | Not written in manuscript                                                                                  | L1 and S2                                                 |
| Rowland et al (2016)10 | No comparator | L1: Pulse width: 200–530 µs, Frequency: 20–40 Hz, Amplitude: 575–650 µA; L2: Pulse width: 300 ms, Frequency: 20–40 Hz, Amplitude: 750 mV | 2 leads were implanted; no details given about company or stimulator | L1 and L2                                                 |

(Continued)
| Study                              | Comparator | Parameters of stimulation                                                                 | Stimulator used (electrode and device)                                                                 | Position of the leads |
|-----------------------------------|------------|------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------|----------------------|
| **Peripheral neuropathy including diabetic peripheral neuropathy**               |            |                                                                                          |                                                                                                         |                      |
| Eldabe et al (2018)               | No comparator | Based on patients' feedback, stimulation was programmed for either subperception or paraesthesia with participants able to adjust parameters using a wireless controller | Up to 4 quadrupolar leads and fully implantable neurostimulation system (Abbott Laboratories)             | L2 and L5           |
| Maino et al (2017)                | No comparator | Pulse width: 670 µs Frequency: 40 Hz Amplitude: 1.88 mA                                    | Quadrupolar DRG lead (Axium, Spinal Modulation, Inc)                                                   | L5                  |
| van Velsen et al (2018)           | No comparator | Not written in manuscript                                                                 | 4-contact Axium” lead (St. Jude Medical, Plano, TX, USA)                                               | L5 and S1           |
| van Velsen et al (2018)           | No comparator | Not written in manuscript                                                                 | Bilateral leads, 4 leads in total (4-contact Axium lead, St Jude Medical, St Paul, MN, USA), stimulator not specified | L5 and S1           |
| **Chronic testicular pain**       |            |                                                                                          |                                                                                                         |                      |
| Hassanain and Murphy (2018)       | No comparator | Pulse width: 130 µs Frequency: 20 Hz Amplitude: 0.55–0.6 mA                               | Proclaim DRG Implantable Pulse Generator IPG (Abbott Laboratories) and Axium Neurostimulator System Slim Tip A50 leads (Abbott Laboratories) | L1                  |
| **Anterior cutaneous nerve entrapment syndrome**                               |            |                                                                                          |                                                                                                         |                      |
| Mol and Roumen (2018)             | No comparator | Not written in manuscript                                                                 | DRG Axium Neurostimulator                                                                                | T9–T12 and L2       |
| **Loin pain hematuria syndrome**  |            |                                                                                          |                                                                                                         |                      |
| Zuidema and Schapendonk (2017)    | No comparator | T12: Pulse width: 300, 130, 180 ms Frequency: 20 and 24 Hz Amplitude: 0.7, 0.95, and 0.775 mA L1: Pulse width: 300, 130, 140, and 180 ms Frequency: 20 and 24 Hz Amplitude: 0.18, 0.6, 0.7, and 0.2 mA | Axium permanent stimulator and quadrupolar stimulation lead (Spinal Modulation, Inc)                   | T12 and L1          |

**Abbreviations:** CRPS, complex regional pain syndrome; DRG, dorsal root ganglion.

fied as active or recruiting with no results (details are given in Table S3).

Results for efficacy and selection of participants for different pain syndromes

**Complex regional pain syndrome**

CRPS was the most common indication treated with EFS neuromodulation of DRG among studies included in this review, with nine such studies (Table 1). ACCURATE, an RCT published in 2017, included 152 participants and compared neuromodulation with EFS of DRG with traditional SCS.1 The remaining studies had nonrandomized designs. Three studies were before and after comparisons, including a total of 71 participants during the trial period of stimulation and 44 for permanent implantation.2,3,31 There were five case reports that included 17 participants in total.32–36
The ACCURATE trial by Deer et al included participants suffering from chronic, intractable pain for at least 6 months, who have tried and failed at least two prior pharmacologic treatments from two different drug classes. The trial results showed that the proportion of participants who achieved treatment success at 3 months in DRG EFS neuromodulation group of participants (81%) was statistically higher compared to the group treated with SCS (56%). A similar result was observed at 12 months follow-up when 74% of participants in DRG EFS neuromodulation group and 53% in SCS group still had significant pain relief. For both the follow-up time points, the results demonstrated DRG stimulation statistical noninferiority ($P<0.0001$) but also statistical superiority ($P<0.0004$). DRG stimulation also demonstrated greater improvements in quality of life and psychological disposition when compared to SCS. Beside CRPS, 32 participants in DRG arm and 33 in SCS arm were diagnosed with causalgia. At 3 months follow-up, when results were stratified by primary diagnosis, higher proportion of participants from DRG arm (79.3%) met the primary endpoint, in comparison with SCS arm (53.3%).

Two uncontrolled before–after comparisons from Liem et al included participants with chronic intractable pain who have failed other treatment modalities (pharmacological and/or surgical) and followed them for 6 months and 12 months after a permanent stimulator implantation. Results showed pain reduction by 66.1% from baseline immediately after implantation of permanent stimulator and remained stable at 6 ($P<0.001$) and 12 months ($P<0.005$) follow-up visit, with 56.1% reduction from baseline. These studies also measured the psychological aspects of pain management using 30-item Brief Profile of Mood States (POMS) and showed statistically significant improvement in four out of six domains of the POMS, as well as decrease in the total mood disturbance score. Third before–after study by van Bussel et al in 2018 compared the efficacy of dorsal column (DC) stimulation vs DRG EFS neuromodulation. Participants had a trial period of 16 days with two stimulation types, SCS and DRG. Reduction of pain was comparable between two groups, but most of the participants preferred DRG stimulation ($P=0.04$) since they did not feel stimulation-induced paresthesia and did not have to adjust stimulation intensity during the day, which was necessary for SCS stimulation.

Among the five case reports, Yang et al (2017) reported 2 cases of implantation of DRG stimulation system after the failure of traditional SCS, in which both the participants reported sustained pain relief at 8 months follow-up. A case report of van Bussel et al (2015) included one participant, followed for 3 months; major pain relief was reported after 8 days, 1 and 3 months. The remaining three case reports included a total of 17 participants who have previously failed various treatment; they measured pain intensity by numeric rating scale (NRS); and all participants showed >50% pain reduction at all follow-up time points (Table 1).

Analysis of participants with CRPS included in studies warrants division of those studies in two groups, where one group included participants in an RCT or an observational study with clear inclusion and exclusion criteria and another group included participants reported in a case series or case report without such criteria.

In studies with defined inclusion and exclusion criteria, common criteria for inclusion were that participants were aged above 18 years, diagnosed with CRPS for at least 6 months, had visual analog scale (VAS) scores of at least 50 or 60 mm out of 100 mm, had failed previous treatment including pharmacological and surgical, were naive to stimulation, had stable neurologic function, and were free from psychological pathology that contraindicated an implantable device (Table 2). Exclusion criteria were participants who already had an implantable device, had previously failed SCS therapy, had cognitive, physical, or sensory impairment, had a coagulation disorder or uses anticoagulants, and if pregnant or planning pregnancy (Table 2). On the contrary, in all case studies and case reports, DRG stimulation was the last treatment option when participants failed all other treatment modalities including also, in some cases, SCS (Table 2). Generally, participants included in these nine studies were older than 50 years, and only one study reported race of participants, with >90% of white participants included. Therefore, the results of these studies are not necessarily generalizable of the population suffering from CRPS.
studies had fixed parameters with pulse width between 160 µs and 362 ms, frequency between 20 and 46 Hz, and amplitude between 500 and 1,030 µA. Details of neuromodulation with EFS parameters used in analyzed studies are given in Table 3.

**Low back pain**

Four uncontrolled before–after studies included 33 participants with LBP. Deer et al (2013) included 10 participants with pain of different etiologies (including peripheral neuropathy and postherpetic neuralgia), but most of the participants had LBP; so we included it in this group (Table 2).³⁸ The average reduction in pain between baseline and final visit was 70%±32% (P=0.0007). Time of last follow-up for different participants varied from 6 months up to 2 years.³⁸

Uncontrolled before–after studies of Huygen et al and Weiner et al included participants with LBP after failed back surgery syndrome (FBSS). Huygen et al (2018) found that at 12 months follow-up, VAS was reduced by 44.2% (P<0.001) from baseline.⁶ The other study of Huygen et al included participants with multiple etiologies; 25 participants with LBP due the FBSS, 13 diagnosed with causalgia, 11 with CRPS, and few participants with several others etiologies. From 56 included participants, 49 who were implanted with permanent stimulation system were followed up to 12 months, 49% of them reported ≥50% pain relief, whereas 82% reported at least 30% reduction from baseline (P<0.0001).⁴⁰ Both the studies reported sustained improvements in mood measured by POMS.⁶,⁴⁰ In Weiner et al (2016), after 6 weeks >50% reduction in VAS was achieved in 63% of participants.³⁷ Parameters of stimulation and details about stimulator used are given in Table 3.

Two studies in this group were case reports that used a novel high-frequency type of neuromodulation with EFS with parameters of stimulation different from standard DRG stimulation. One was case report including only one patient with fixed stimulation parameters that showed 66% improvement for back pain and 56% for leg pain at 6 months follow-up.³ The other was a small feasibility study that included six participants and used a range of stimulation parameters, specifically amplitude between 1 and 24 mA, pulse width of 10–1000 µs, and frequency of 2–10,000 Hz. The study reported that all participants achieved >50% pain reduction from baseline to 3 months follow-up (Table 3).³⁹

Main inclusion criteria for participants in this group of studies were at least 18 years old, diagnosed chronic pain syndrome, VAS scores ≥5 or 6 out of 10, failure of other treatment modalities. Participants were excluded if they had an active implantable stimulator of any type, if pregnant, and if they had any inability to comply with study requirements (Table 2).

**Groin pain**

Groin pain was analyzed in one uncontrolled before–after comparison with 34 participants,⁴¹ one case series with 25 participants suffering from groin pain of different etiology⁴⁰ and one case report describing three participants.⁷ Morgalla et al had follow-up for as much as 3 years and showed significant decrease in pain after 3 months, after 1 and 2 years (P=0.001), and after 3 years (P=0.005), when compared to the baseline measurement.⁴¹ Schu et al reported >80% reduction in VAS in 47.8% and >50% reduction in VAS in 82.6% of participants for an average follow-up time period of 27.8±4.3 months.⁸ Zuidema et al also reported significant reduction in VAS scores at 2 and 3 months follow-up (Table 1).⁷

Participants included in studies about groin pain were all aged ≥18 years, diagnosed with chronic groin pain, and have failed previous treatment modalities (Table 2). Participants from case report had also failed treatments with transcutaneous electrical nerve stimulation (TENS) and pulsed radiofrequency prior to successful treatment with DRG stimulation.⁷ Detailed parameters of stimulation used in those studies are given in Table 3.

**Pelvic girdle pain**

Two case reports with eight participants analyzed effect of neuromodulation with EFS of DRG on pelvic girdle pain.⁹,¹⁰ Hunter et al reported significant pain relief in all seven participants after a trial implantation period, whereas four participants reported sustained pain relief 1 year after permanent implantation.⁹ Rowland et al reported a case of 43% pain reduction from baseline using NRS after 6 months.¹⁰ Included participants had failed various treatments including medication, neurolysis, surgery, steroid injections, and, in some cases, SCS (Table 2). Details about stimulation parameters are given in Table 3.

**Peripheral neuropathy and diabetic peripheral neuropathy**

One case series included ten participants with peripheral diabetic neuropathy who had an average VAS pain reduction from baseline by 64.2%±35.8% (P<0.001) at 12 months.⁴³ Participants with peripheral neuropathy were also described in three case reports, of which two reported the same patient, and hence we analyzed only once.²⁹,³⁰,⁴² In both cases, participants reported >50% pain relief after 12 months follow-up. Included participants have failed other treatment modalities.
including use of neuropathic pain medications. In van Velsen et al\textsuperscript{39} and Eldabe et al\textsuperscript{43} published in 2018, participants also tried traditional SCS, whereas in Maino et al\textsuperscript{42} TENS was applied prior to neuromodulation with EFS of DRG (Table 2). Only study by Maino et al\textsuperscript{42} reported small fiber neuropathy diagnosed by skin biopsy, which confirmed pathological reduction of intraepidermal unmyelinated nerve fibers. Other included studies did not clearly document the type of peripheral neuropathy.

Other chronic pain states represented in a single study
This category includes participants with various painful conditions represented by single case series or case report with less than ten participants. One case report included eight participants with phantom limb pain who were treated with DRG stimulation after a failure of other treatment modalities. The percentage of pain reduction was on average 52\%±31.9\% from baseline during an average follow-up time period of 9 months.\textsuperscript{44} Other case report included five participants with ACNES. Three participants had good pain relief after 12 months follow-up, whereas two were refractory to the therapy without any pain relief.\textsuperscript{47}

Other pain conditions treated with DRG stimulation were chronic intractable pain in the coccyx with 90\% pain reduction at 4 months,\textsuperscript{45} chronic testicular pain with sustained pain reduction of 70\%–80\% during 1 year,\textsuperscript{46} and LPHS with >50\% pain relief after 3 years.\textsuperscript{48} These case reports included only one participant.

Details about efficacy and safety of treatment for those indications are given in Table 1, included participants in Table 2 and about parameters of stimulation in Table 3.

Results about safety
Results about safety of neuromodulation with EFS of DRG could be classified as related to the procedure, related to the device, or related to the stimulation technique. In the group of SAEs related to the procedure, the most common event was infection at the site of implantation. SAEs related to the device included infection of stimulator pocket site, dural puncture, postdural puncture headache, and transient loss of function. The most common SAE related to stimulation was overstimulation. One participant died 6 months after implantation due the medication overdose. This was attributed to previously existing depression.\textsuperscript{40} Several other SAEs occurred that could not be classified as related to implantation procedure or stimulation including depression, bladder infection, bowel obstruction, pain following a capsaicin (Qutenza) application, perianal fistula, knee cyst, transient ischemic attack, worsening of pre-existing CRPS, and temporary loss of leg strength. AEs included loss of stimulation, leads migration, pain at incision site, and postprocedure headache. Incidence of AEs for each study is shown in Table 1.

High proportion of case reports did not report any safety data, ten from included 29 studies. While several studies explicitly mentioned that no complications occurred, it remained unclear if that was true for those studies that did not mention AEs at all.

RoB assessment
RoB in included studies was assessed using Cochrane RoB tool for the one RCT that was included. We judged domains for random sequence generation and allocation concealment as unclear RoB, as those methods were not reported. Blinding of both participants/personnel and outcome assessors was judged with high RoB because the study was not blinded. The risk of attrition bias was judged as low, since authors reported all attrition during trial, as well as during follow-up period and performed modified intention-to-treat analysis when reporting results. From included 76 participants in both arms, 61 in DRG arm and 54 in SCS arm completed trial period, whereas 12 months follow-up was completed by 55 participants in DRG arm and 50 participants in SCS arm.

Reasons for exclusion or failure of treatment were given. We considered that the study had unclear risk of selective reporting bias because in the registered protocol only primary outcome was mentioned and secondary outcomes shown in the manuscript were not mentioned in the protocol. We did not find other sources of bias (Table S4).

Attempts to conduct meta-analysis
In our study protocol, we planned to conduct a meta-analysis of outcomes reported in RCTs. However, we were unable to conduct a meta-analysis since we found only one RCT.\textsuperscript{1}

Reporting of conclusion statements for efficacy and safety in manuscript abstracts
We were also interested in determining the proportion of studies that reported conclusion statements about efficacy and safety in manuscript abstracts, since sometimes abstracts are the only source of information for clinicians. Such statements for efficacy were either positive conclusive (N=12) or positive inconclusive (N=13). The remaining four studies did not report conclusion statements about efficacy (Table S5).
In the majority of included studies, the abstract did not include any conclusions about the safety of a tested intervention (N=19). In the remaining abstracts, there were positive conclusive (N=5) or positive inconclusive statements about safety (N=3), whereas two abstracts only provided information about number of AEs or mentioned certain specific AEs, without providing overall conclusions about safety (Table S5).

Funding and conflict of interest in included studies
Most of the included studies had conflict of interest statements. In almost 60% of the studies, authors reported that they either are consultants of companies providing financial support to the research or that they have equity in those companies. Only nine studies reported that authors have no conflict of interest.

Overall, among 29 studies, there were eight industry-funded studies, only two studies were financed by a nonprofit institution, and none were funded by government or other grant sources. In the group of industry-funded studies, four out of eight had positive conclusive statements about intervention, while four had positive inconclusive statements. Among 21 non-industry funded studies, we found eight studies with positive conclusive statements and nine with positive inconclusive statements. However, a majority of studies that mentioned a potential conflict of interest did not explicitly mention sources of funding, so we were unable to judge whether those were funded by industry or are they more likely to yield positive findings about intervention.

Discussion
This systematic review included 29 small studies about the use of EFS of DRG as neuromodulation method for treatment of pain. We found that studies about neuromodulation with EFS of DRG reported participants treated for painful conditions of various etiologies, but mostly in participants who have failed many or all other available treatment modalities. For some participants it was reported that they were refractory to stimulation and that they did not experience any pain relief. The majority of studies that reported conclusion statements about efficacy in their abstracts indicated that there is positive, but inconclusive evidence regarding efficacy of neuromodulation with EFS of DRG. We were unable to perform meta-analysis since only one of the 29 included studies was RCT.

Several reviews have been published recently on this topic, but with a narrower focus and number of methodological limitations. Harrison et al (2017) published a literature review about efficacy and safety of DRG stimulation as a treatment for neuropathic pain. Chang Chien et al (2017) published a systematic review about alternate intraspinal targets for SCS. This review covered very wide range of topics, and DRG stimulation was just one of the analyzed interventions. The review searched only the single database PubMed, whereas the Cochrane Handbook and CRD guidelines indicate that a systematic review requires a search of at least two bibliographic databases. These reviews had also several additional methodological limitations. Specifically, the authors used very simple search strategies, some of them did not report the search dates, they did not report excluded studies, there were no analyses of RoB in included studies, attempts to make quantitative analysis were not reported, and potentially competing interests of authors of included studies were also not reported. None of the studies were focused on participant selection or reported parameters of stimulation.

The latest review published by NACC had some elements of systematic review methodology, including search strategy and analysis of quality of included studies, and gave very comprehensive overview of the topic with sections on DRG anatomy and physiology, with the main focus on DRG stimulation devices and implantation procedure. However, authors mentioned participants’ selection very briefly, reported parameters of stimulation only for ACCURATE study and had last date of search in June 2017, which is currently >1 year ago. In addition, consensus evidence and given recommendations were partly based on published abstracts without inclusion of full manuscripts.

From all chronic painful conditions treated with neuromodulation with EFS of DRG, CRPS was the condition with evidence represented by ten of the 29 studies included in this systematic review and also with greatest overall number of participants included. Furthermore, this was only condition for which evidence about efficacy and safety was available from RCT. We rated RoB as unclear for multiple domains due to the lack of information provided in manuscript. NACC used modified Pain Physician criteria and US Preventive Services Task Force (USPSTF) criteria to give final grading. The group rated ACCURATE study as level 2 according to modified Pain Physician criteria and level I using USPSTF criteria and overall recommended DRG stimulation as an effective therapy for treatment of CRPS type I of the lower extremity, while for upper extremity CRPS type I or II conclusion was that more studies are needed.

NACC had also strong consensus about the use of DRG stimulation for groin pain, rating the overall evidence as

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NACC had also strong consensus about the use of DRG stimulation for groin pain, rating the overall evidence as
level II-2. This is in agreement with our findings since, together with LBP, those were painful conditions represented with uncontrolled before-after studies of higher quality and including more participants than case series and case reports, which reported results for the rest of included chronic painful symptoms. Further studies are needed with higher level evidence about efficacy and safety of neuromodulation with EFS of DRG for treatment of those conditions.

We excluded studies published only as conference abstracts, as it has been shown that such information are not necessarily dependable, as authors may change results, either qualitatively or quantitatively, by the time data from conference abstracts are published in peer-reviewed journals. Since our search is dated September 2018, we included several painful conditions that have been treated with DRG EFS neuromodulation for the first time such as chronic intractable pain in the coccyx, ACNES, and LPHS. We also included small pilot studies and case reports that used novel high-frequency parameters of stimulation.

This evidence is of low quality, represented only with few participants to whom neuromodulation with EFS of DRG was the last treatment option after failure of other treatment modalities. Median number of participants in these studies was 6. Based on these findings, future larger studies should also consider inclusion of participants diagnosed with these conditions to confirm safety and efficacy of therapy, as well as use of novel stimulation parameters which could possibly improve treatment outcomes.

Weaknesses of available evidence

We found that a source of funding was not reported in the majority of included studies. Most of the studies that had financial support were funded by industry with commercial interest in neuromodulation with EFS of DRG, which warrants cautious interpretation of the results. Furthermore, studies that reported potential conflicts of interest in which the authors were either consultants of the industry producing the studied device or had equity in those companies did not report the source of funding. A recently published systematic review about industry sponsorship and research outcome in studies of drugs and medical devices found that industry-sponsored research more frequently reported favorable efficacy results and favorable conclusions for tested intervention compared to studies with nonprofit funding. The authors did not find any difference for the majority of RoB items between commercially and nonprofit-funded studies, suggesting that existence of “industry bias” cannot be explained by standard RoB domains. Amiri et al (2014) reported similar results by analyzing >1,300 studies in the field of spine research. They found significant associations between source of funding, study outcome, and level of evidence, in which unfunded and industry-funded studies had the highest proportion of level IV evidence and reported a higher proportion of favorable outcomes, while studies with public funding or funding other than industry had a higher proportion of level I evidence.

Since the source of funding may influence outcomes, reporting sources of funding and conflicts of interest should be a mandatory part of each manuscript. Researchers should follow ethical principles and transparency when reporting study results, while clinicians should critically appraise each paper they are reading, not relying exclusively on the authors’ conclusions.

We were aware of the limited number of RCTs in the field as well as the fact that other study designs had lower methodological quality, less reliable results, and thus provide lower level of evidence about certain treatments. However, NRSDs can be valuable sources of information, having longer follow-up time, especially regarding safety of intervention, which was our outcome of interest, so we decided to include also NRSDs in this systematic review.

Even though we followed criteria for conducting a high-quality systematic review, our evidence synthesis has limitations that are related to the published studies on this topic. The evidence is based on studies with small number of participants, whereas there was only one RCT, and a large proportion of case series and case reports. More reliable evidence is needed to make reliable conclusion about efficacy and safety of studied intervention.

Conclusion

EFS of DRG is a widely used neuromodulation intervention for treating various painful conditions of different etiologies. Studies published thus far imply that the intervention may help highly selected participants with various pain syndromes, who have failed to achieve adequate pain relief with other pharmacological and nonpharmacological interventions. Some participants were refractory to the treatment, without any pain relief. However, these findings need to be taken with extreme caution because of multiple limitations of available studies. These limitations include poor quality of available studies, very small number of participants included, highly selected patient population who participated in these studies, and conflict of interest of sponsors and authors of those studies. Due to availability of only one trial on this topic, with high or unclear RoB on the majority of analyzed domains, currently available evidence from studies...
on humans about benefits of neuromodulation with EFS of DRG for treatment of pain should be considered preliminary and confirmed in high-quality RCTs with sufficient number of participants.

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Disclosure
The authors report no conflicts of interest in this work.

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