

# The effect of HF10™ Therapy on opioid analgesic use for the treatment of chronic intractable low back and/or radicular leg pain

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		<input type="checkbox"/> Protocol
<b>Registration date</b> 16/10/2018	<b>Overall study status</b> Completed	<input type="checkbox"/> Statistical analysis plan
		<input type="checkbox"/> Results
<b>Last Edited</b> 17/12/2019	<b>Condition category</b> Musculoskeletal Diseases	<input type="checkbox"/> Individual participant data
		<input type="checkbox"/> Record updated in last year

## Plain English summary of protocol

### Background and study aims:

Long-term opioid use is common in people with chronic pain. This can lead to immune suppression, endocrine suppression, reduced libido, and an overall detrimental effect on the quality of life. These side effects have pushed the need for more appropriate pain management options, such as spinal cord stimulation (SCS). SCS involves implanting a device that delivers small electrical fields to the spinal cord to provide pain relief in patients with chronic (long-term), intractable (hard to control) back and leg pain. Significant leg pain relief has been reported with traditional SCS which uses a low frequency (<1200 Hz), but there is limited long-term data supporting the use of low frequency SCS in predominant back pain patients. Recent studies testing SCS using a frequency of 10 kHz (HF10 Therapy™) have shown that this treatment is effective and safe in patients with both back and leg pain. The results from these studies demonstrated that these patients had significant pain relief up to 24 months. The decreased pain in both back and leg were consistent throughout the studies with improvements in functional capacity with no perception of paresthesia (a burning or prickling sensation). This study will investigate the effect of HF10 Therapy on opioid analgesic pain management in a low back pain and/or leg pain population.

### Who can participate?

Adults with chronic intractable low back and/or radicular leg pain

### What does the study involve?

Participants undergo Spinal Cord Stimulation (SCS) at 10 kHz (HF10 Therapy) as per the center's usual practice. SCS involves the surgical placement of two leads (which look like very thin wires) into a small area near the spinal cord. Electrical stimulation is delivered through these wires by a small, battery-operated, rechargeable SCS implanted generator. Each patient is followed for 12 months after device activation. Following device activation, participants will consult with their doctor to actively work towards reducing their pain medications during the 12 month follow up period. The participants attend regular clinic visits to complete questionnaires, and to provide feedback on their pain. Successfully treated participants get a permanent device implanted as per center's routine clinical practice.

What are the possible benefits and risks of participating?

Participants may benefit from the SCS system to relieve their pain. There are no direct risks of taking part in this study, although the general risks of having a spinal stimulation device in place apply.

Where is the study run from?

7 pain centers across Australia

When is the study starting and how long is it expected to run for?

January 2018 to December 2022

Who is funding the study?

Nevro Corp (USA)

Who is the main contact?

Mr Wim Laloo

laloo@nevro.com

## Contact information

### Type(s)

Scientific

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Public

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## Additional identifiers

### Protocol serial number

CA2018AU OR

# Study information

## Scientific Title

A Prospective Post-market Study to Investigate the Effect of HF10™ Therapy on Opioid Analgesic Use for the Treatment of Chronic Intractable Low Back and/or Radicular Leg Pain

## Study objectives

To investigate the effect of HF10 Therapy on opioid analgesic pain management in chronic intractable low back and/or radicular leg pain population.

## Ethics approval required

Old ethics approval format

## Ethics approval(s)

Bellberry Limited, 17/09/2018, ref 2018-08-633

## Study design

Observational prospective multi-centre cohort study

## Primary study design

Observational

## Study type(s)

Treatment

## Health condition(s) or problem(s) studied

Chronic intractable low back and/or radicular leg pain

## Interventions

As part of their standard care, participants undergo a trial (temporary evaluation period) of Spinal Cord Stimulation (SCS) at 10 kHz (HF10 Therapy) as per the centre's usual practice and following this, those for whom the trial treatment was successful have a permanent device implanted as per center's routine clinical practice.

Following device activation, participants are followed for a period of 12 months. Following device activation, participants will consult with their doctor to actively work towards reducing their pain medications during the 12 month follow up period. The participants attend regular clinic visits, at 1, 3, 6, 9 and 12 months, to complete questionnaires, and to provide feedback on their pain, quality of life, disability, health status, satisfaction and sleep. At these visits, the physicians will take assessments from the participants such as pain evaluation, medication usage and side effects (if any).

## Intervention Type

Device

## Primary outcome(s)

The proportion of subjects who report clinical success with HF10 therapy at the 6-month study visit. Clinical success is denoted by any of the following criteria described on the Opioid Use and Pain Outcome Matrix:

1. Decreased opioids ( $\geq 30\%$ )
2. Decrease in predominant pain ( $\geq 30\%$ )

### **Key secondary outcome(s)**

The following will be successively evaluated (hierarchical test approach) in the order shown with a 0.05 significance until statistical significance is not achieved:

1. Proportion of subjects who respond to HF10 therapy, assessed at the baseline and after 6 months by a  $\geq 50\%$  improvement in low back pain or leg pain visual analogue scale (VAS) from the baseline at 6 months
2. Proportion of subjects reporting at least a 30% reduction from baseline in opioid analgesic use as measured by the morphine equivalent daily dose (MEDD) at 6 months
3. Proportion of subjects who respond to HF10 therapy, assessed at the baseline and after 12 months by a  $\geq 50\%$  improvement in low back pain or leg pain VAS from the baseline at 12 months
4. Proportion of subjects reporting at least a 30% reduction from baseline in opioid analgesic use as measured by the morphine equivalent daily dose (MEDD) at 12 months
5. Proportion of subjects that experience at least a 10-point reduction in the Oswestry Disability Index at 12 months
6. Proportion of patients reporting  $\geq 0.100$  improvement from the baseline in quality of life, assessed using the EQ-5D-5L at 12 months

### **Completion date**

31/12/2022

## **Eligibility**

### **Key inclusion criteria**

1. Diagnosed with chronic intractable back and/or leg pain
2. 18 years of age or older at time of enrolment
3. Currently taking opioids within a dose range of 15-100 mg/day oral morphine or the equivalent (MEDD)
4. Average back or leg pain intensity of  $\geq 5$  out of 10 cm on the Visual Analog Scale (VAS) at enrolment
5. Meet the requirements for being an HF10 Therapy trial candidate as per the centers' practice, which includes psychological evaluation
6. Willing and able to complete health questionnaires and pain scales as specified in the protocol

### **Participant type(s)**

Patient

### **Healthy volunteers allowed**

No

### **Age group**

Adult

### **Lower age limit**

18 years

### **Sex**

All

### **Key exclusion criteria**

1. Plans to enrol in another clinical study during their participation in this study, or are currently enrolled in an interventional clinical study that could interfere in participation in this study or affect the scientific soundness of this study
2. Medical condition or pain in other area(s), not intended to be treated with SCS, that could interfere with study procedures, accurate pain reporting, and/or confound evaluation of study endpoints, as determined by the Investigator (such as primary headache diagnosis or fibromyalgia)
3. Current addiction to cocaine, opiates, alcohol, or benzodiazepines as determined by the treating physician
4. Evidence of an active disruptive psychological or psychiatric disorder or other known condition significant enough to impact perception of pain, compliance of intervention and/or ability to evaluate treatment outcome, as determined by either the treating physician or a psychologist
5. Current diagnosis of a progressive neurological disease such as multiple sclerosis, chronic inflammatory demyelinating polyneuropathy, rapidly progressive arachnoiditis, rapidly progressive diabetic peripheral neuropathy, brain or spinal cord tumor, and/or central deafferentation syndrome
6. Current diagnosis of a coagulation disorder, bleeding diathesis that would put subject at any increased risk of bleeding during SCS procedure, progressive peripheral vascular disease, or uncontrolled diabetes mellitus
7. Diagnosis of scoliosis that precludes lead placement
8. Condition currently requiring or likely to require the use of diathermy
9. Metastatic malignant disease or active local malignant disease
10. Life expectancy of less than 1 year
11. Active systemic or local infection
12. Pregnant (if female and sexually active, subject must be using a reliable form of birth control, be surgically sterile or be at least 2 years post-menopausal)

**Date of first enrolment**

31/12/2018

**Date of final enrolment**

31/12/2021

## **Locations**

**Countries of recruitment**

Australia

**Study participating centre**

**Northern Integrated Pain Management**

Suite 3, 20 Smith Street,

Charlestown

Australia

NSW 2290

**Study participating centre**  
**Interventus Pain Specialists**  
Suite 20A Level 10  
Evan Thomson Building  
Chasely Street  
AUCHENFLOWER,  
Australia  
QLD 4066

**Study participating centre**  
**Pain Management Unit**  
The Canberra Hospital,  
Building 8, Level 1,  
Off Palmer Street Garran,  
Woden  
Australia  
ACT 2606

**Study participating centre**  
**Precision Brain and Spine**  
115 Cotham Road  
Kew  
Australia  
VIC 3101

## **Sponsor information**

**Organisation**  
Nevro Corp

**ROR**  
<https://ror.org/02xcxe208>

## **Funder(s)**

**Funder type**  
Industry

**Funder Name**  
Nevro Corp

# Results and Publications

## **Individual participant data (IPD) sharing plan**

The datasets generated during and/or analysed during the current study are/will be available upon request from Dr Adele Barnard

## **IPD sharing plan summary**

Available on request