



### Interventional Pain Management: Opioid-Sparing Technologies

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### Disclosure

- Consultant/Independent Contractor: Abbott, Avanos, Biotras, Nalu, SI-Bone, Nevro, Vertos Medical, Vertiflex/Boston Scientific
- Grant/Research Support: Avanos, Biotronik, Sollis Pharmaceutical, Semnur Pharmaceutical, Nevro, Vertiflex
- Advisory Board: Biotras, Nalu, Nevro, Vertiflex



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### Learning Objectives

- Review pain and analgesia
- Discuss the impact of chronic pain
- Describe the evolution of opioid therapy
- Review current and future application of technology in treating chronic pain
- Review supporting evidence



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### Outline

- Chronic pain
- History of analgesia
- Evolution of pain opioid therapy
- Technologies in treating chronic pain
  - Neuromodulation
  - Minimally invasive spinal interventions
- Evidence review in opioid reduction
- Explore the latest clinical trials



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### Pain

- "An unpleasant sensory and emotional experience associated with actual or potential tissue damage, or described in terms of such damage..."



PainWeek

Merskey H, Bogduk N et al. IASP Task Force on Taxonomy, 1994

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"Like a rope ringing a bell"



FIG. 1-1. Descartes' (1664) concept of the pain pathway. He writes: "If for example fire (A) comes near the foot (B), the minute particles of this fire, which as you know move with great velocity, have the power to set in motion the spot of the skin of the foot which they touch, and by this means pulling upon the delicate thread (C) which is attached to the spot of the skin, they open up at the same instant the pore (d-e) against which the delicate thread ends, just as by pulling at one end of a rope makes to strike at the same instant a bell which hangs at the other end." From Melzack, R., and Wall, P.D.: Pain mechanisms: A new theory. Science, 165(371), 1965.

PainWeek

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### Origin of Analgesia



- Sumerians, 3000 B.C. who first cultivated the poppy plant for its opium
- Homer in 300 B.C. Helen of Troy to treat her grief over the absence of Odysseus
- Morphine, Codeine, Heroin, Oxycodone

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### Ancient Pain Management



Auricular acupuncture depicted during Han dynasty, 200 BC



Cauterizing the external ear to treat migraine, 12<sup>th</sup> century Persian surgery text

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### Morphine



- Discovered by Friedrich Serturmer in 1803
- Named after Morpheus, the god of dreams
- Commercially made available by Merck in 1827

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### Opioid Problem is Not New

- 1849, Mrs. Charlotte Winslow, Bangor, Maine
- 65 mg morphine per ounce
- "sooth any human or animal...effectively quieted restless infants and small children, especially for teething"



**Painweek** [https://en.wikipedia.org/wiki/Mrs.\\_Winslow%27s\\_Soothing\\_Syrup](https://en.wikipedia.org/wiki/Mrs._Winslow%27s_Soothing_Syrup)

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### Diacetylmorphine



- Alder Wright, 1874 by adding 2 additional acetyl groups
- 4x more potent than morphine
- Manufactured by Bayer
- Prescribed in the U.K. for withdrawal and analgesic
- Schedule I substance in U.S.

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### Contemporary Anesthesia



- Oct. 16, 1846, William Morton demonstrates the use of ether for dental extraction at Massachusetts General Hospital
- Surgeon, John Warren,
- "Gentleman , this is no humbug."

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### Chronic Pain in America

- 1 in 5 Americans suffer from chronic pain
- Large economic impact: ~\$600 billion/year
- Loss of productivity: ~\$300 billion/year
- Opioid epidemic: #1 health crisis in America
- National health survey by NIH 2012
  - 50 million adults experience pain every day
  - Pain → worse overall health status
  - Female, elderly, non-Hispanics (Asians less likely)



A Certified Tool to Improve Care for Seriously Ill Hospitalized Patients.  
[www.painweek.com/related-topics/trends-statistics/overdose-death-rates](http://www.painweek.com/related-topics/trends-statistics/overdose-death-rates)

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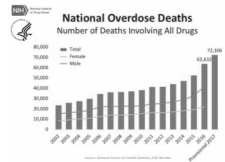
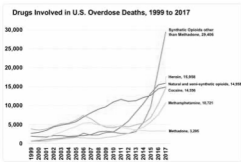
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### Opioid Crisis in America



- Over 72,000 Americans died in 2017 from drug overdose
- More than 49,000 deaths involved opioids
- Synthetic opioid deaths have surged



<https://www.drugabuse.gov/related-topics/trends-statistics/overdose-death-rates>

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### Paradigm Shift in Opioid Therapy

- Lack of long term efficacy for treating chronic pain
- Risk for tolerance, dependency, and abuse
- National opioid crisis
- New CDC opioid prescribing guidelines



<https://www.cdc.gov/drugoverdose/prescribing/guidelines.html>

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### Evolution of Pain Medicine

In contrast to earlier thinking on the order of treatments in the pain treatment continuum,<sup>1</sup> it has been proposed that device therapies be considered at an earlier stage.<sup>2</sup>

\*Kramka ES. Intrathecal Opioid Therapy for Nonmalignant Pain. Current Perspectives 2015 Clinical Guidelines. J Pain Symptom Manage. 2015;50(1):1-11.  
 †Schnitzler, DR. et al. Live Your Life Pain Free. October 2005. Based on the interventional pain management expertise of Dr. John Schoenrock.

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### Emergence of Electroceuticals

- Bioelectronics
- Therapeutic devices
- External or implanted
- Delivering electricity
- Neuromodulation
- Alter disease states
- Market prediction of \$35.5 billion global market by 2025

1. Kristoffer Frum, Nature, 2013  
 2. <http://www.grandviewresearch.com/press-release/global-electroceuticals-bioelectric-medicine-market>

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### Ancient Opioid-Sparing Technologies

- Baghdad Battery
- 250 BC, outside Baghdad
- Clay jar with asphalt stopper
- Iron rod surrounded by copper
- If filled with vinegar: 1.1 volts
- Torpedo fish
- 46 AD: Scribonius Largus used torpedo fish to treat chronic pain

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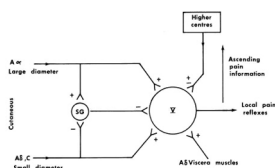
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### Gate Theory of Pain



- Wall and Melzack, 1965
- Aβ (sensory) and Aδ, C pain fibers compete for passage through physiologic "gate"
- Stimulation of larger Aβ fibers would: closes the gate



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### 50 Years of Spinal Cord Stimulation

Electrical Inhibition of Pain  
By Stimulation of the Dorsal Columns:  
*Preliminary Clinical Report*

**I**ntermittent electrical stimulation of the dorsal columns...  
REPORT OF A CASE  
 A 50-year-old male patient with severe, chronic, intractable pain...  
REFERENCES



Dr. Shealy



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### Contemporary Landmark Studies

- Kemler, et al. NEJM. 2000
  - SCS vs. PT alone in treatment of CRPS (n=54)
  - at 6 mo. 58% of SCS compared to 6% of PT improved
- North, et al. Neurosurgery. 2005
  - Re-operation vs. SCS with crossover (n=50)
  - 47% SCS vs. 12% re-op improved
  - 37% crossover, and 43% achieved pain relief
- Manca, et al. PROCESS Trial, Eur. J. Pain. 2008
  - SCS vs. CMM for FBSS
  - SCS with improved health and function, but higher \$
- Kumar, et al. Neurosurgery. 2008
  - SCS vs. CMM alone for 6 month with crossover (n=100)
  - at 24 mo. 37% of SCS compared to 2% CMM



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### Spinal Cord Stimulation



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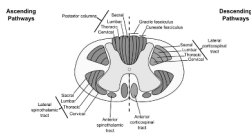
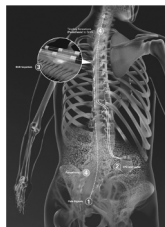
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### Traditional SCS Therapy



- Electrical stimulation of dorsal column
- Activation of Aβ sensory fibers
- Generate paresthesia in areas of pain

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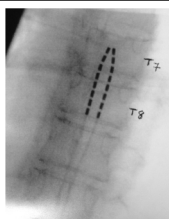
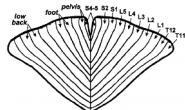
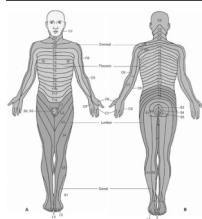
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### Paresthesia Dependent SCS Therapy



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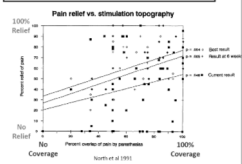
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### Paresthesia Dependent SCS Therapy

Traditional SCS Paradigm:  
More paresthesia overlap = more pain relief



- Paresthesia coverage of pain is considered necessary for efficacy
- Paresthesia mapping
- Advanced lead placement

**PainWeek** North et al 1991

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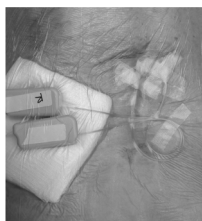
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### SCS Trial



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### Renaissance of Neuromodulation



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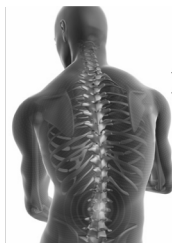
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### Innovations in Neuromodulation

- Adaptive stimulation
- MRI compatibility
- Novel wave forms
- Novel targets of stimulation
- Closed loop technology
- Vagal nerve stimulation



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### Adaptive Stimulation

- To address intensity variations due to postural changes
- Distance to spinal cord changes with posture
- Accelerometer controlled programming options
- 41% reported reduction of daily adjustments<sup>1</sup>
- First use of feed back in SCS



PainWeek

J. Schultz, et al. Pain Physician, 2012

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### Novel Targets of Stimulation

- Dorsal root ganglion
- Vagal nerve stimulation
- Peripheral nerve stimulation
- Multifidus stimulation



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### Paresthesia Free Stimulation

- "High Density": ~ 1kHz, top of the traditional "low frequency" range, adjusted below perceptual threshold
- "High Frequency": 10 kHz, beyond perceptual threshold
- "Burst": 500 Hz x 5 pulses x 40/sec, totaling 200/sec, adjusted below perceptual threshold
- Differential Targeted Multiplexed (DTM) wave forms to target multiple cell types



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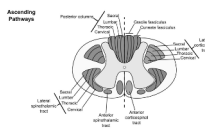
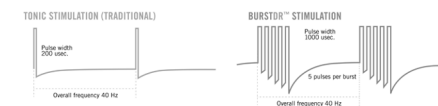
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### Burst Waveform in SCS Therapy



- Target medial descending pathway
- Both pain intensity and quality
- Via C-fiber activation in lamina I
  - Medial thalamic nuclei
  - Anterior cingulate cortex

Expert Review of Medical Devices, 2018

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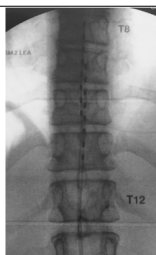
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### High Frequency SCS Therapy



**Comparison of 10-kHz High-Frequency and Traditional Low-Frequency Spinal Cord Stimulation for the Treatment of Chronic Back and Leg Pain: 24-Week Results From a Multicenter, Randomized, Controlled Trial**

**OBJECTIVE:** The aim of this study was to compare the effectiveness of 10-kHz high-frequency spinal cord stimulation (SCS) with traditional low-frequency SCS (LF-SCS) for the treatment of chronic back and leg pain. The study was a multicenter, randomized, controlled trial. The primary endpoint was the mean change in the Visual Analog Scale (VAS) score for back pain at 24 weeks. The secondary endpoints were the mean change in the VAS score for leg pain, the mean change in the Oswestry Disability Index (ODI) score, and the mean change in the EuroQol-5D (EQ-5D) score. The study was conducted in 10 centers across the United States. The study population consisted of 100 patients with chronic back and leg pain. The patients were randomized to receive either 10-kHz high-frequency SCS or LF-SCS. The study was conducted over a 24-week period. The results of the study showed that 10-kHz high-frequency SCS was significantly more effective than LF-SCS for the treatment of chronic back and leg pain. The mean change in the VAS score for back pain was significantly greater in the 10-kHz group compared to the LF-SCS group. The mean change in the VAS score for leg pain, the mean change in the ODI score, and the mean change in the EQ-5D score were also significantly greater in the 10-kHz group compared to the LF-SCS group. The study was limited by its short duration and the lack of a placebo control group. The study also had a high rate of dropouts. The study was funded by the National Institutes of Health.



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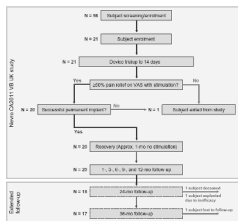
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### Al-Kaisy NSRBP Pilot Study Design



Single Arm, Prospective Study

- 20 successful implants
- 3 year observation
- Predominant back pain
  - Baseline 7.9cm VAS
- Multiple outcomes assessed:
  - Opioid usage
  - Function (ODI)

Published results at 12 and 36 months



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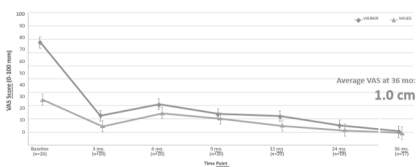
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### Non-Surgical Back Pain Pilot Study: 36 Months

Non-Surgical Back Pain Pilot – 36 Months  
Now Published in Pain Medicine



1. Al-Kaisy, Ahmad, Pankaj, Siddiqui, Smith, Thomas E, Caviglioli, Raj, Bhargava, Michael, Peng, David, Bergman, William, Lam, Aksh, Lavin, Jonathan. Long-Term Improvement in Chronic Axial Low Back Pain Patients With Percutaneous Spinal Surgery: A Cohort Analysis of NSRBP. High-Frequency Spinal Cord Stimulation over 36 Months. Pain Medicine 2017; 18: 1-8.  
2. Bopp, Richard G. Future surgery for lumbar degenerative disc disease: still more questions than answers. The Spine Journal 15 (2004): 272-274.



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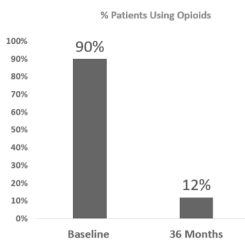
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### NSBP Study: Significant Reduction in Opioids

- 90% of patients on opioids at baseline
- 12% of all subjects were using opioids at 36 months



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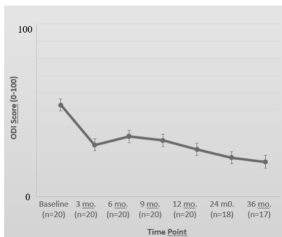
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### NSBP Study: Significant Improvement of Function

Average ODI of 53 at baseline  
 - "Severe Disability"

Average ODI of 19.8 at 36 months  
 - "Minimal disability"



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### Dorsal Root Ganglion SCS Therapy



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### PAIN

OPEN

#### Dorsal root ganglion stimulation yielded higher treatment success rate for complex regional pain syndrome and causalgia at 3 and 12 months: a randomized comparative trial

Timothy R. Deer<sup>1\*</sup>, Robert M. Levy<sup>2</sup>, Jeffrey Kramer<sup>3</sup>, Lawrence Poree<sup>4</sup>, Kasra Amirjalilian<sup>5</sup>, Eric Griggby<sup>6</sup>, Peter Stastik<sup>7</sup>, Alan W. Burton<sup>8</sup>, Abram H. Burchiel<sup>9</sup>, Jon Olson<sup>10</sup>, James Scowcroft<sup>11</sup>, Stan Gobovac<sup>12</sup>, Leonardo Kuper<sup>13</sup>, Richard Pascau<sup>14</sup>, Christopher Kent<sup>15</sup>, Jason Poppe<sup>16</sup>, Thomas Yearwood<sup>17</sup>, Sam Samuel<sup>18</sup>, W. Porter McRoberts<sup>19</sup>, Homer Cassini<sup>20</sup>, Mark Netherton<sup>21</sup>, Nathan Miller<sup>22</sup>, Michael Schaudt<sup>23</sup>, Edward Tavel<sup>24</sup>, Timothy Davis<sup>25</sup>, Kristina Davis<sup>26</sup>, Linda Johnson<sup>27</sup>, Nagy Mekhalif<sup>28</sup>

- U.S. pivotal trial, comparing DRG and traditional stimulation
- Multi-center, randomized controlled trial
- 152 subjects with CRPS, causalgia of the lower extremity
- 76 DRG, 76 SCS
- At 3 months DRG group 81.2% and SCS group 55.7% efficacy



Deer T. et al. Pain, 2017

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### Recent Landmark Studies

- Accurate Trial: pivotal U.S. study DRG stimulation
- Sunburst Trial: pivotal U.S. study for Burst
- SENZA RCT: pivotal U.S. study for HF10
- Accelerate Trial: HF-SCS versus conventional SCS
- Avalon Trial: closed loop SCS study in Australia
- Evoke Trial: pivotal U.S. study for closed loop SCS
- Acute Trial: pivotal U.S. study for DTM



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### Real World Results

High-Volume Centers Study Shows Real World Outcomes Comparable to SENZA-RCT

#### Design

- 1660 consecutive patients enrolled (2014-2018)
- Eight global, high-volume HF10 centers

#### Long Term Efficacy (n=1100\*)

- 78% responder rates
  - 74% responder rates in prior SCS patients
- 90% satisfaction
- 32% of patients reduced medication intake
- 3.7% reported explant rate
  - 1.2% due to loss of efficacy



Rizzo, Thomas et al. A Multicenter Real-World Review of HF10 SCS Outcomes for Treatment of Chronic Back &/or Limb Pain. *Annals of Clinical and Translational Neurology*. January 2019. See only the 2176 patients who safety data available. Of the 2176 patients, 1176 (54%) were enrolled in the SENZA RCT, 1000 (46%) were not. Of the 1000 patients, 1100 (11%) were enrolled in the SENZA RCT, 890 (89%) were not. The mean time between implantation and the last visit was 6.9 months (range 0.1-33.2).

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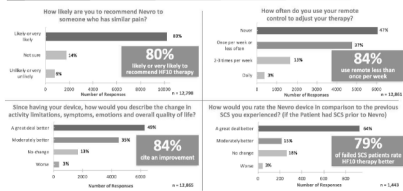
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### HF10 SCS: My Results

#### COMMERCIAL PATIENT FOLLOW UP – TEXT & CALL RESULTS



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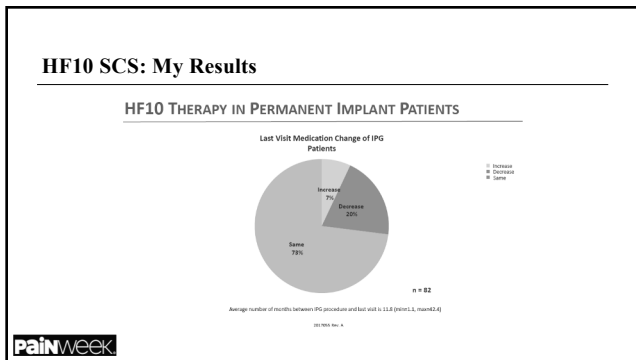
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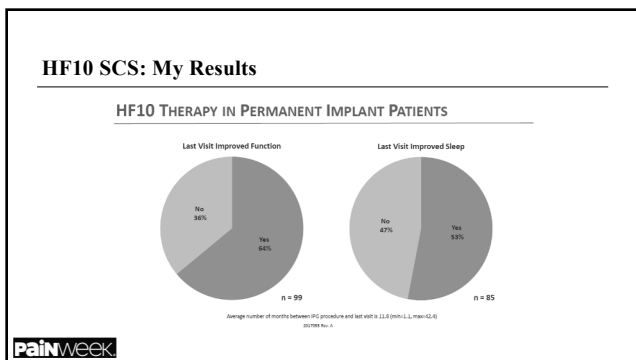
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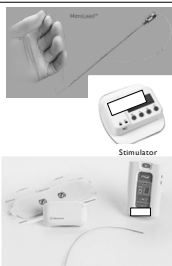
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### PNS for Chronic and Acute Pain

- FDA approved
- 0.2mm coiled lead via 20g introducer needle
- Coiled lead design for tissue ingrowth
- Temporary and revisable
- External wearable power source
- Forgiving lead placement
- Low infection risk



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### PNS for Chronic and Acute Pain

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#### One Year Follow-Up of a Randomized, Double-Blind, Placebo-Controlled Trial of Percutaneous Peripheral Nerve Stimulation for Chronic Pain Following Amputation

Rosenfarb JM, MD<sup>1</sup>, Gilmore CA, MD<sup>2</sup>, Iqbal BM, MD<sup>3</sup>, Li S, MD<sup>4</sup>, Desai MJ, MD<sup>5</sup>, Hunter CW, MD<sup>6</sup>, Rauck RL, MD<sup>7</sup>, Naderi A, MD<sup>8</sup>, Mahr J, MD<sup>9</sup>, Cohen SP<sup>10</sup>, Crosby ND, PhD<sup>11</sup>, Boggs AH, PhD<sup>12</sup>  
<sup>1</sup>Department of Neurological Surgery, Northwestern University, Chicago, IL, USA; <sup>2</sup>Center for Chronic Research, Winston-Salem, NC, USA; <sup>3</sup>Department of Anesthesiology, University of California San Diego, San Diego, CA, USA; <sup>4</sup>Division of Pain Management, The Ohio State University, Columbus, OH, USA; <sup>5</sup>Department of Neurology, University of Michigan, Ann Arbor, MI, USA; <sup>6</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>7</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>8</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>9</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>10</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>11</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>12</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA.

**INTRODUCTION**

MINIMALLY INVASIVE PERCUTANEOUS PNS  
 • Percutaneous peripheral nerve stimulation (PNS) has been shown to be effective for the treatment of chronic pain following amputation.  
 • This study evaluated the efficacy of PNS in a randomized, double-blind, placebo-controlled trial.

**MATERIALS & METHODS**

**Study 1 (PNS Treatment)**  
 • Randomized, double-blind, placebo-controlled trial.  
 • Participants: 40 patients with chronic pain following amputation.  
 • Interventions: PNS (n=20) vs. Placebo (n=20).  
 • Outcomes: Pain intensity, functional status, and quality of life.

**Study 2 (PNS Treatment)**  
 • Randomized, double-blind, placebo-controlled trial.  
 • Participants: 40 patients with chronic pain following amputation.  
 • Interventions: PNS (n=20) vs. Placebo (n=20).  
 • Outcomes: Pain intensity, functional status, and quality of life.

**RESULTS**

**Acute Pain Intensity**  
 • 87% of the treated group (PNS) and 60% of the control group (Placebo) reported a decrease in acute pain intensity at 12 weeks.

**Chronic Pain Intensity**  
 • 87% of the treated group (PNS) and 60% of the control group (Placebo) reported a decrease in chronic pain intensity at 12 weeks.

**Functional Status**  
 • 75% of the treated group (PNS) and 50% of the control group (Placebo) reported an improvement in functional status at 12 weeks.

**DISCUSSION**

Percutaneous PNS treatment over 12 weeks may provide significant and enduring pain relief for patients with chronic pain following amputation.

**REFERENCES**

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#### Reductions in Opioid Consumption with Percutaneous Peripheral Nerve Stimulation (PNS) of the Lumbar Medial Branch Nerves for Chronic Low Back Pain

Christopher Gilmore, MD<sup>1</sup>, Leonardo Kapural, MD, PhD<sup>2</sup>, Thomas Hopkins MD, MBA<sup>3</sup>, Mehul Desai, MD, MPH<sup>4</sup>, Michael DePalma, MD<sup>5</sup>, Sean Li, MD<sup>6</sup>, Abram Burdick, MD<sup>7</sup>, Timothy Deer, MD<sup>8</sup>, Meredith McGee, PhD<sup>9</sup>, Joseph Boggs, PhD<sup>10</sup>  
<sup>1</sup>Center for Chronic Research, The Ohio State University, Columbus, OH, USA; <sup>2</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>3</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>4</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>5</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>6</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>7</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>8</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>9</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA; <sup>10</sup>Department of Neurology, University of Colorado Denver, Denver, CO, USA.

**INTRODUCTION**

MINIMALLY INVASIVE PERCUTANEOUS PNS  
 • Percutaneous peripheral nerve stimulation (PNS) has been shown to be effective for the treatment of chronic low back pain.  
 • This study evaluated the efficacy of PNS in a randomized, double-blind, placebo-controlled trial.

**MATERIALS & METHODS**

**Study 1 (PNS Treatment)**  
 • Randomized, double-blind, placebo-controlled trial.  
 • Participants: 40 patients with chronic low back pain.  
 • Interventions: PNS (n=20) vs. Placebo (n=20).  
 • Outcomes: Pain intensity, functional status, and opioid consumption.

**Study 2 (PNS Treatment)**  
 • Randomized, double-blind, placebo-controlled trial.  
 • Participants: 40 patients with chronic low back pain.  
 • Interventions: PNS (n=20) vs. Placebo (n=20).  
 • Outcomes: Pain intensity, functional status, and opioid consumption.

**RESULTS**

**Substantial Reductions in Opioid Analgesic Consumption with PNS**  
 • Significant reductions in opioid consumption were observed in the PNS group compared to the placebo group.

**Chronic Pain Intensity**  
 • Significant reductions in chronic pain intensity were observed in the PNS group compared to the placebo group.

**Functional Status**  
 • Significant improvements in functional status were observed in the PNS group compared to the placebo group.

**DISCUSSION**

Percutaneous PNS treatment over 12 weeks may provide significant and enduring pain relief for patients with chronic low back pain.

**REFERENCES & ACKNOWLEDGEMENT**

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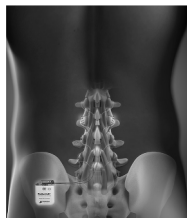
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### Multifidus Stimulation

- Multifidus stimulation
- ReActiv8 Clinical Trial
- N=53, multicentered RCT
- Improvement of chronic LBP
- 58% responder rate at 12 months
- Just published in Neuromodulation



**PainWeek** NeuroModulation, 2018

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### Spinal Stimulation for the Treatment of Intractable Spine and Limb Pain

A Systematic Review of RCTs and Meta-analysis

Tim J. Larmer, MD<sup>1</sup>, Susan M. Moeschler, MD<sup>1</sup>, Helena M. Gazzeika, MD<sup>1</sup>, W. Michael Hooters, MD<sup>1</sup>, Markus A. Bendel, MD<sup>1</sup>, M. Hassan Murad, MD<sup>2</sup>

DOI: <https://doi.org/10.1016/j.mayocp.2018.12.037>



- Systematic review, 12 studies, 980 patients from 1995-2017
- Compare SCS to medical therapy
- SCS increased odds of pain reduction by 50% or more in 3 trials
- SCS significantly reduced VAS in 3 trials
- HF10, Burst, and DRG increased odds compared to traditional SCS

**PainWeek** Larmer T, et al. Mayo Clinic Proceedings, 2019

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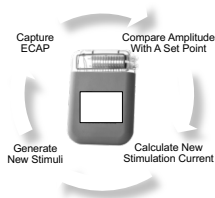
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### Closed-Loop Stimulation

- Not FDA approved
- Measure the response of Aβ fibres to stimulation
- Capture ECAP and make real time adjustments to stimulation
- 1,000,000 times per day
- Maintain stim within individual therapeutic window



**PainWeek**

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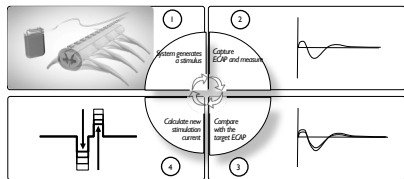
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### Variable Output Feedback Controlled Stimulation



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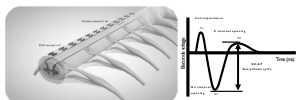
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### What is an ECAP?

- **E**voked **C**ompound **A**ction **P**otentials (ECAPs) are the sum of the electrophysiological response from multiple nerve fibers
- ECAPs provide insight into the type of fibers stimulated and are a measure of spinal cord (SC) activation



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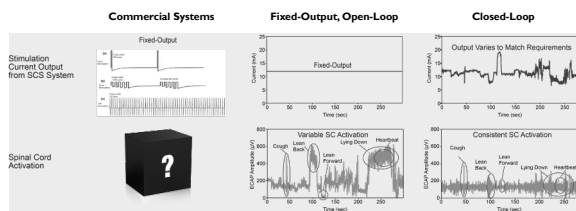
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### Fixed-Output versus Closed-Loop SCS



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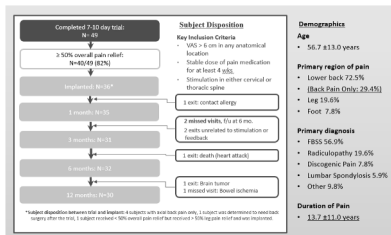
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### Avalon Study (Australia)



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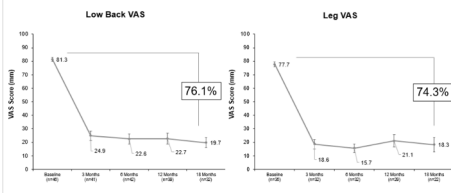
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### Avalon Study Results

Avalon 18-Mos Pain Reduction



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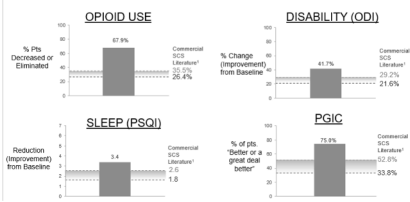
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### Avalon Study Results

Avalon 18-Mos. Secondary Results, Compared to Literature



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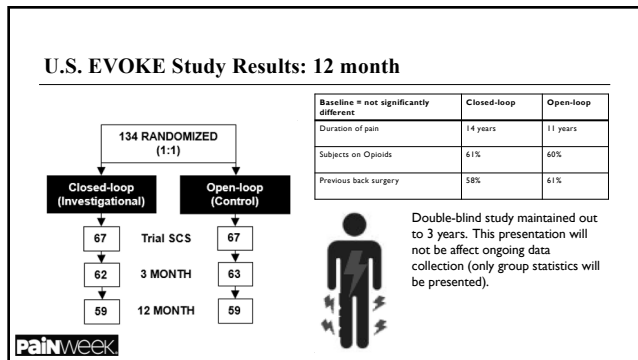
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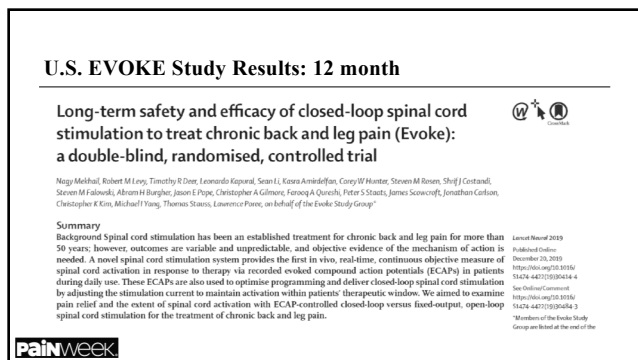
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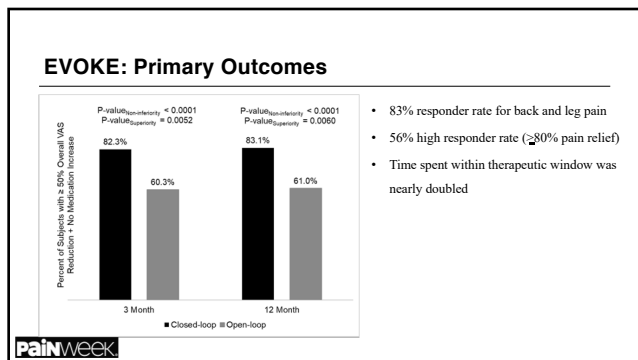
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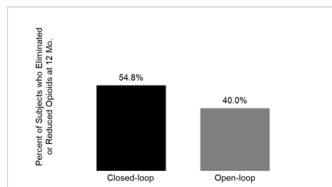
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### EVOKE: Reduction/Elimination of Opioids



MME	Closed-Loop	Open-Loop
Baseline	80	64
12 Months	45	45



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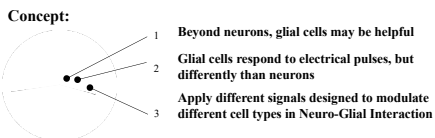
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### Differential Targeted Multiplexed SCS



- Differential Target** Differential target: Different pulse signals intended for different cell types
- Multiplexed** Multiplexed: Multiple pulse signals combined within the delivered stimulation



- Pulse signals within 20-1,200 Hz range & max pulse width of 1 ms
- Multiple programs are applied according to algorithm

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### What Do We Know About Glial Cells?

Maintain a balanced homeostatic state with neurons.  
 Disruptions of the Neuro-Glial Interaction can result in chronic neuropathic pain.

Greatly outnumber neurons in the cord tissue exposed to SCS

# 12:1



Yalçın et al (JNAN 2019)

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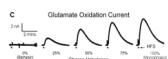
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### Modulating Neuro-Glial Interaction

- Both neurons and different types of glial cells are important to chronic pain (Neuro-Glial Interaction)
- Glial cells are electrically excitable, yet differently than neurons

For example, depending on stimulation patterns, glial cells will release different levels of glutamate



PainWeek

Agarwal et al., J. Neuro. Eng. 2019

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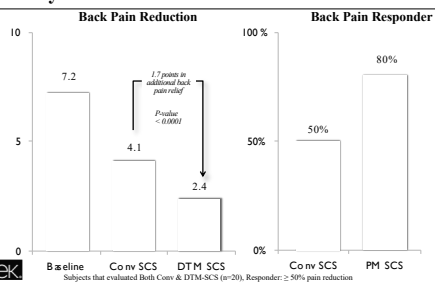
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### Acute Study Results: Back Pain



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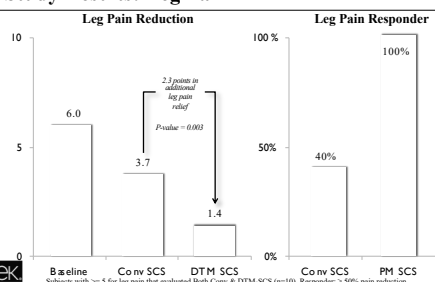
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### Acute Study Results: Leg Pain



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**Ultra Minimally Invasive SCS**

- **Battery-free, microstimulator**
  - Smallest IPG available (<1.5 cc, battery-free)
  - Minimally invasive
  - Potential to decrease rates of pocket pain and infection
- **Small size, without compromise**
  - Highly capable & easily upgradeable
    - Robust connectivity – Confirmation of connection and therapy delivery
    - Multiple therapy options
    - Upgradeable without the need for surgery
- **Potential to expand your practice / patient population**
  - Multiple indications – US Clearance for both SCS & PNS
  - Potential to increase patient acceptance
  - Smartphone app remote control



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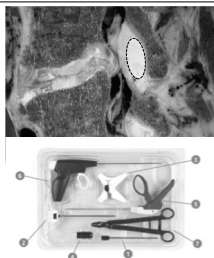
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**LSS Treatment: Percutaneous Image-Guided Decompression (PILD)**

- Debulk the hypertrophied dorsal ligamentum flavum
- Image-guided percutaneous approach
- Key safety factor is the epidurogram
- Ligament greater than 2.5mm
- Outpatient procedure
- Under mild sedation
- 24 month data, MiDAS ENCORE Trial
- Re-Approved by Medicare, 2018



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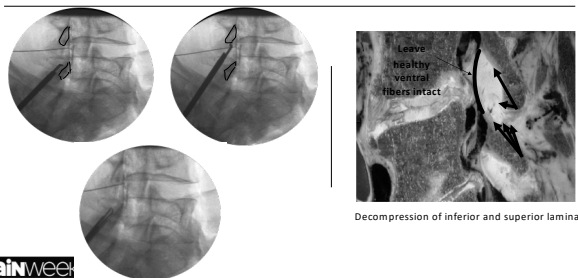
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**LSS Treatment: PILD Procedure**



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### ENCORE Study 2-year Outcomes Confirmed Long-term Safety and Efficacy<sup>3</sup>

#### Study Protocol

- Coverage with evidence development (CED)
- Prospective, multicenter, randomized controlled
- Randomization:
  - mid versus ESI
- Study visits:
  - Baseline, 6 month, 1 year, 2 years
- Comparative data through 1 year
  - mid-only at 2 years
- Outcome measures:
  - Oswestry Disability Index (ODI)
  - Numeric Pain Rating Scale (NPRS)

#### Study Population

- Patients experiencing neurogenic claudication symptoms
- Hypertrophic ligamentum flavum
  - > 2.5 mm
- 65 years or older
- ODI > 31
- NPRS > 5
- No surgery at any treatment level
- Spondylolisthesis
  - < Grade III

**PainWeek** Shoup PL, Chaffin TB, Calverley S, et al. Long-term safety and efficacy of minimally invasive lumbar decompression procedure for the treatment of lumbar spinal stenosis with neurogenic claudication: 2-year results of MIDAS ENCORE. Reg Anesth Pain Med. 2018;43:789-794.

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### ENCORE Study 2-year Outcomes Functional and Pain Improvement Compared to ESIs<sup>3</sup>

#### Oswestry Disability Index (ODI)

Significant and sustained functional improvement through 2-year follow-up

Mean ODI improvement of 22.7 points at 2 years (10 point improvement is clinically significant.)

#### Numeric Pain Rating Scale (NPRS)

Significant and durable reduction of pain through 2-year follow-up

Mean NPRS improvement of 3.6 points at 2 years (2-point improvement is clinically significant.)

**PainWeek** Shoup PL, Chaffin TB, Calverley S, et al. Long-term safety and efficacy of minimally invasive lumbar decompression procedure for the treatment of lumbar spinal stenosis with neurogenic claudication: 2-year results of MIDAS ENCORE. Reg Anesth Pain Med. 2018;43:789-794.

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### ENCORE Study 2-year Outcomes Significant Improvement by Stenosis Type<sup>3</sup>

#### Stenosis Type: Percent of Patients

Majority of patients had multiple types of stenosis

#### ODI Mean Point Change

Significant functional improvement regardless of stenosis type

**PainWeek** Shoup PL, Chaffin TB, Calverley S, et al. Long-term safety and efficacy of minimally invasive lumbar decompression procedure for the treatment of lumbar spinal stenosis with neurogenic claudication: 2-year results of MIDAS ENCORE. Reg Anesth Pain Med. 2018;43:789-794.

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**LSS Treatment: Interspinous Process Decompression (IPD)**

- Various spacers have been introduced
- Superion is the only percutaneous device
- Serves as a back stop preventing compression of the spinal canal and lateral recess during extension



**Painweek**

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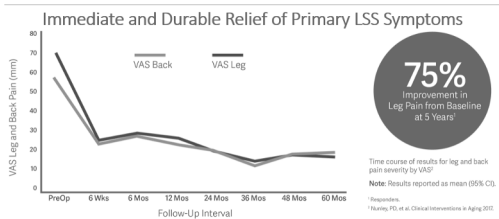
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**LSS Treatment: IPD 5 Year IDE Study Results**



**Painweek**

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Journal of Pain Research

Dovepress

CLINICAL TRIAL REPORT

**Interspinous process decompression is associated with a reduction in opioid analgesia in patients with lumbar spinal stenosis**

- **85%** reduction in the proportion of subjects using opioids at 5 years
- Interspinous process decompression is associated with decrease in the need for opioid medications

**Painweek**

Nancy, PD et al. J Pain Research, 2018

78

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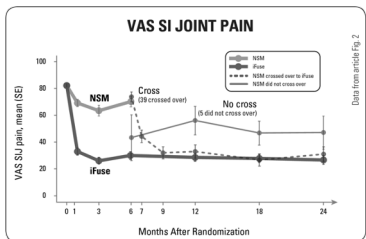
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**INSITE 2-year Results: VAS SI Joint Pain  
Improves more after SI joint fusion than NSM**



*PainWeek - Int J Spine Surg 2016 (INSITE 2yr)*

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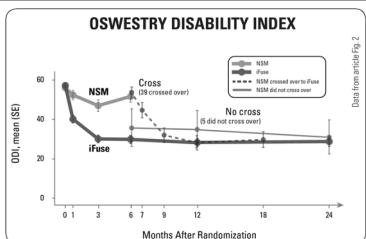
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**INSITE 2-year Results: ODI  
Improves more after SI joint fusion than NSM**



*PainWeek - Int J Spine Surg 2016 (INSITE 2yr)*

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**INSITE 2-year Results**

		iFuse % subjects	NSM % subjects
<b>Primary Endpoint*</b>	Success @ 6 mo	82%	26%
<b>Patient Satisfaction</b>	Very or somewhat satisfied	90% (6 mo)	61% (6 mo)
		88% (2 yr)	
<b>Clinical Improvement</b> (Minimum Clinically Important Difference)	VAS improvement ≥ 20pt	83% (2 yr)	10% (2 yr)
	ODI improvement ≥ 15pt	68% (2 yr)	7.5% (2 yr)
<b>Opioid Use</b>	% change in number of subjects taking opioids	30% ↓ (baseline to 2 yr)	7.5% ↑ (baseline to 6 mo)

\* Binary success/failure composite measure. Success if all criteria met: VAS SI joint pain reduction ≥ 20 points, no device-related SAEs, no neurological worsening, and no surgical re-intervention for SI joint pain.



*PainWeek - Int J Spine Surg 2016 (INSITE 2yr)*

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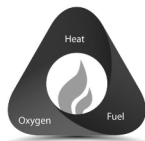
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### Summary

- Opioid Epidemic
  - Unmet treatment needs
  - Health economics
- Chronic pain
  - #1 cause of disability
  - Aging population
- Innovation
  - Technology
  - Level I evidence



Future of interventional pain management is bright

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### Questions

In addition to greater than 50% relief in pain and reduction of VAS score, several interventional pain procedures have show level I evidence for opioid reduction. They include:

- a. Percutaneous sacroiliac joint fusion
- b. High frequency spinal cord stimulation
- c. Interspinous process decompression
- d. Closed loop spinal cord stimulation
- e. All of the above (correct answer)




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### Questions

Various clinical trials in interventional pain management are now incorporating metrics other than pain scores such as the VAS. Additional clinical study end points include:

- a. Functional status in the form of disability index (ODI)
- b. Sleep (PSQI)
- c. Opioid reduction
- d. Severity of neurogenic claudication (ZCQ)
- e. All of the above (correct answer)




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**Questions**

A 75 year old female presents with chronic back and leg pain due to multi-level degenerative disc disease. She has tried various conservative treatment options such as physical therapy, acupuncture, anti-inflammatories, and anti-convulsants. Patient has consulted with a spine surgeon who did not think she was an ideal surgical candidate. In addition to long-term opioid therapy, what other interventional pain therapy should she be considered for?

- a. Interspinous process decompression
- b. Sacroiliac joint fusion
- c. High frequency spinal cord stimulation (correct answer)
- d. Peripheral nerve stimulation
- e. Percutaneous image-guided decompression



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**Thank You**



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