

NeurAxis, Inc.



Reimagining an Evidence-Based, Drug Free Alternative For Children

August 27, 2024

NeurAxis is committed to providing solutions that create value and provide better patient outcomes. We believe in improving lives and minimizing suffering. Through innovation and research, we are reimagining the future of patient care.

Forward Looking Statements

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This PowerPoint discusses ongoing research activities and investor-directed information with percutaneous electrical nerve field stimulator (PENFS) technology. Please note, that the PowerPoint includes information about technology and intended uses of that technology which have not been reviewed or approved/cleared by the U.S. FDA, and is being provided for informational purposes only. NeurAxis does not recommend or suggest the use of its PENFS™ IB-Stim™ device for uses beyond those that are cleared by the U.S. FDA. See https://ibstim.com/important-information/.



NeurAxis PENFS¹: First FDA Indicated Treatment for Pediatric FAP/IBS²



Strong Data = Strong Policy Coverage & Reimbursement = Strong Revenue Growth

Large Global Market with Significant Unmet Need

- \$23B+ TAM³ for target pipeline indications
- \$9B+ TAM³ for target pediatric indications (near-to-mid term)
- \$14B+ TAM³ for target adult indications (mid term)
- Large unmet clinical need: high refractory, off label pharmacological treatments with adverse side effects

Clear Commercial Pathway

- FDA De Novo clearance
- Technology specific CPT billing code
- Major Insurance Payer Coverage initiated
- Strong IP on Device and Method









Unique, Innovative Product Supported by Clinical Evidence

- Novel treatment targeting the brain-gut-axis
- Differentiated PENFS technology
- 700+ published patients⁴
- Easy-to-learn and efficient procedure

Seasoned Management and Board

- Experienced management team and Board of Directors
- Operations and infrastructure built to scale
- Path to profitability



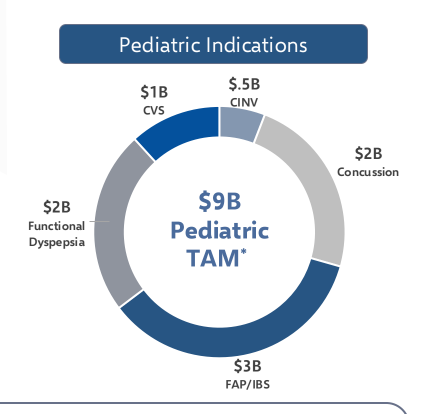
^{1.} Percutaneous Electrical Nerve Field Stimulation

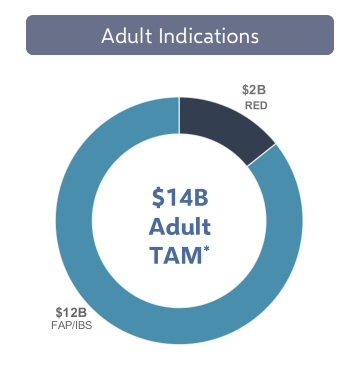
^{2.} FAB/IBS: Functional Abdominal Pain/Irritable Bowel Syndrome

^{3.} Total Addressable Market (TAM) - Calculated by the total number of patients we target to treat multiplied by the revenue potential from each patient

^{4.} Published patient - a patient who went through a study and the study was analyzed and now the study has been published in a peer-reviewed journal

\$23B+ Total Addressable U.S. Market for Pipeline Indications





Why Pediatrics?

- Significant unmet need
- Lack of FDA approved treatment options
- Single call point for future indications

Entering Pediatric markets first with:

- First FDA cleared treatment for Pediatric FAP/IBS
- Growing Body of Clinical Evidence
- Coding, Coverage and Payment
- KOL and AAP/NASPGHAN endorsement



DGBIs: A Problem with an Unmet Need

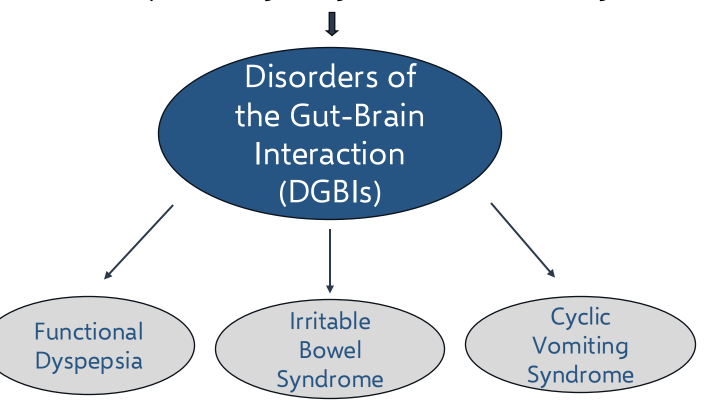
(IB-STIM)

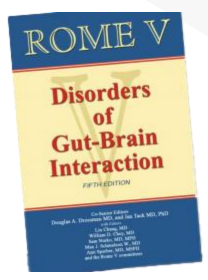
- No FDA-approved therapies for children with abdominal pain-related disorders of the gut-brain interactions (DGBIs)
- Disorders negatively impact quality of life and ability to function (attend school, sports, and social activities)
- Insufficient data to support the use of the most prescribed drugs, some with serious side effects
- A growing number of families and providers are seeking non-pharmacologic alternatives for children



Gut-Brain Axis

Abnormal processing of signals to and from the gut









Functional Gastrointestinal Disorders: Disorders of Gut-Brain Interaction



Functional Dyspepsia – pain or discomfort located in the upper abdomen

Irritable Bowel Syndrome (IBS) –characterized by abdominal discomfort or pain associated with defecation or a change in bowel habit.

Cyclic Vomiting Syndrome – recurrent episodes of intense nausea and vomiting lasting hours to days with intervals of normal wellbeing lasting weeks to months.



Data Does Not Support Standard Pharmacotherapy in Children with IBS



No data to support use of Antidepressants in Children with Functional Abdominal Pain:

- Amitriptyline (TCA) did not beat placebo in RCT¹
- Citalopram (SSRI) did not beat placebo in RCT²

Significant Risk of TCA Side Effects in Children:

- Increased risk of suicidal ideation (black box warning)³
- Mood changes
- EKG disturbance⁴
- Long-term risk of dementia⁵

Substantial Patient Need for Safe & Effective, Non-Pharmacological Alternatives:

 Growing number of families seeking alternative therapies for pain in children⁶



^{1.} Saps M, Youssef N, Miranda A, et al. Multicenter, randomized, placebo-controlled trial of amitriptyline in children with functional gastrointestinal disorders. *Gastroenterology*. 2009;137:1261-1269.

^{2.} Roohafza H, Pourmoghad das Z, Saneian H, Gholamrezaei A. Citalopram for pediatric functional abdominal pain: a randomized, placebo-controlled trial. Neurogastroenterol Motil. 2014;26:1642-1650.

^{3.} Jick H, Kaye JA, Jick SS. Antidepressants and the risk of suicidal behaviors. JAMA. 2004;292:338-343.

^{4.} Chogle A, Saps M. Electrocardiograms changes in children with functional gastrointestinal disorders on low dose amitriptyline. WorldJ Gastroenterol. 2014;20:11321-11325.

Coupland CAC, Hill T, Dening T, Morriss R, Moore M, Hippisley-Cox J. Anticholinergic Drug Exposure and the Risk of Dementia: A Nested Case-Control Study [published online ahead of print, 2019 Jun 24]. JAMA Intern Med. 2019;179:1084-1093.

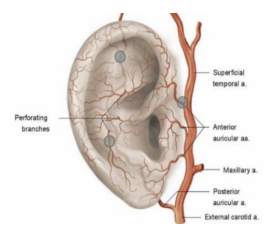
^{6.} Groenewald CB, Beals-Erickson SE, Ralston-Wilson J, Rabbitts JA, Palermo TM. Complementary and Alternative Medicine Use by Children With Pain in the United States. Acad Pediatr. 2017;17:785-793.

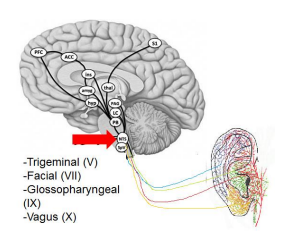
Percutaneous Electrical Nerve Field Stimulation (PENFS)



How Does Neuromodulation Work?







1. Access

Direct access to central nervous system (CNS) through peripheral cranial nerves

2. Stimulate

Stimulation reduces firing of amygdala

3. Change

Induces changes in brain pathways/connectivity



Established Technology with Demonstrated Safety and Efficacy







What is IB-Stim™

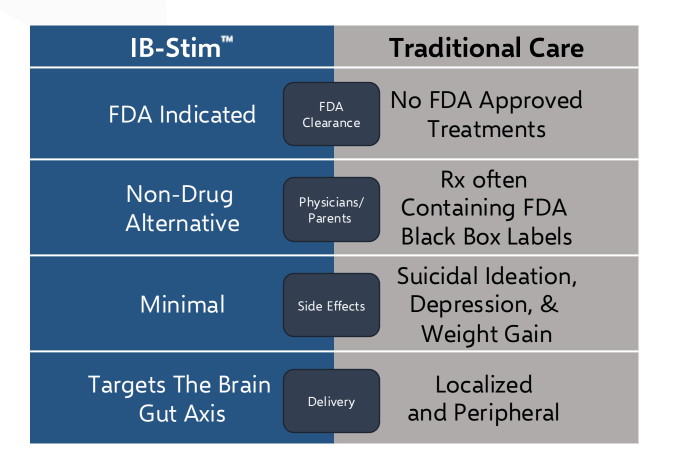
- PENFS system intended for patients 11-18 years of age with functional abdominal pain (FAP) associated with IBS
- Aids in pain reduction via neuromodulation to branches of Cranial Nerves (V,VII,IX and X)
- Non-drug and non-surgical device therapy that can be placed in an outpatient clinic
- Used 120 hours per week for up to 3-4 consecutive weeks*

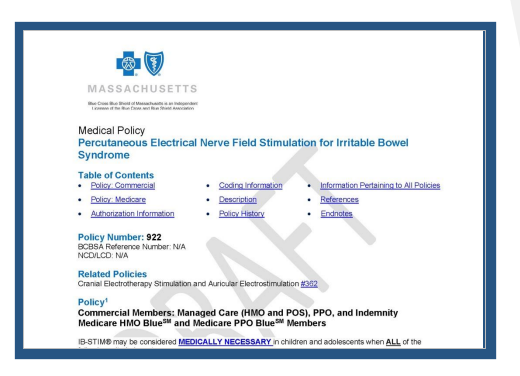
FDA De Novo Clearance

CPT CAT III Effective July 1,2022



IB-Stim™ Advantages Over Traditional Care







Patient Journey





1. Persistent Pain

Patient experiences frequent and often debilitating abdominal pain (weeks, months or years)



5. Follow-Up

- Patient takes off at home after 5 days, gets a 2-day break, then visits a physician for next prescribed treatments for up to 4 weeks.
- Further follow up visits / titration as needed



2. Consultation

General Pediatrician

- Pain is generalized, non-specific, showing no "red flags"
- Counsels on lifestyle changes
- If no benefit, trial of medication
- If no benefit, referral to Pediatric Gastroenterologist

Pediatric Gastroenterologist

- Blood work (CBC, metabolic panel, inflammatory markers, celiac screen) and stool test
- If negative, treatment with medication is started
 - o Antidepressants (TCAs and SSRIs) used for pain
 - Anti-histamine (Cyproheptadine)
- Anti-spasmodics (Hyoscyamine)
- o Cognitive behavioral therapy, where available

Families often skip PCP since referral is not required



FDA cleared IB-Stim™ can be used first vs. traditional, off-label pharmacotherapy-based approach



4. Use/Care

- Stays on for 120 hours (5 days)
- No special care requirements except to avoid getting wet



3. Placement

- Outpatient (in-office) procedure placement by acting Physician
- Requires no anesthesia



IB-Stim™ Research – By the Numbers

Strong Data = Strong Policy Coverage & Reimbursement = Strong Revenue Growth



16 Current Publications
Utilizing NeurAxis' PENFS Technology

10 Types of Studies



Double Blind
Placebo Controlled



Long-Term Data



Registry Data

Clinical fMRI Study



Quality of Life Data



Real World Clinical Data

Animal Mechanistic Study

Head-to-Head vs. SoC

Health Economic Study

Safety Data

13 Children's Hospital Study Sites















IB-Stim Publications in Children with DGBIs

Effect of percutaneous electrical nerve field stimulation on mechanosensitivity, sleep, and psychological comorbidities in adolescents with functional abdominal pain disorders

Neha R Santucci¹ | Christopher King² | Khalil I. El-Chammas¹ Anundorn Wongteerasut¹ | Alisara Damrongmanee¹ | Kahleb Graham¹ | Lin Fei³ | Rashmi Sahay³ | Cheryl Jones¹ | Natoshia R. Cunningham⁴ | Robert C Coghill²

Percutaneous Electrical Nerve Field Stimulation in Children and Adolescents With Functional Dyspepsia—Integrating a Behavioral Intervention

Neha R. Santucci, MD^{1,2}; Alan J. Beigarten, MS¹; Fatima Khalid, MS¹; Khalil I. El-Chammas, MD^{1,2}; Kahleb Graham, MD^{1,2}; Rashmi Sahay, MD³; Lin Fei, PhD3: Kristin Rich, PhD2,4: Michael Mellon, PhD2,

Prospective study of the effect of auricular percutaneous electrical nerve field stimulation on quality of life in children with pain related disorders of gut-brain interaction

Ashish Chogle^{1*}, Kaajal Visnagra², Jamie Janchoi^{1,3}, Tammy Tran¹, Rachel Davis³, Nicole Callas¹ and Elisa Ornelas^{1,3}

I ne micropiome in adolescents with irritable bowel syndrome and changes with percutaneous electrical nerve field stimulation

Nicholas J. Ollberding^{2,5} | Pachmi Sahay⁵ | Naha P. Santucci^{1,2}

Percutaneous Electrical Nerve Field Stimulation for Drug-Refractory **Pediatric Cyclic Vomiting Syndrome**

Karrento. Katia MD*; Venkatesan, Thangam MD[†]; Zhang, Liyun MSc[‡]; Pawela, Louis BS^{*}; Simpson, Pippa PhD[‡]; Li, B U.K. MD^{*}

Percutaneous electrical nerve field stimulation compared to standard medical therapy in adolescents with functional abdominal pain disorders

Neha R. Santucci^{1,2*}, Rashmi Sahay³, Khalil I. El-Chammas^{1,2}, Kahleb Graham¹², Mikaela Wheatlev¹², Madeleine Vandenbrink², Jennifer Hardy¹ and Lin Fei³

Percutaneous Electrical Nerve Field Stimulation Compared to Standard Medical Therapy in Adolescents with Functional Abdominal Pain Disorders

Neha R. Santucci^{1, 2*} Rashmi Sahay¹ Khalil I. El-Chammas^{1, 2} Kahleb Graham^{1, 2} Mikaela Wheatley^{1, 2} Madeleine Vandenbrink¹

Percutaneous Electrical Nerve Field Stimulation Improves Comorbidities in

Katja Karrento^{1*}, Liyun Zhang², William Conley¹, Zeeshan Qazi³, Thangam Venkatesan⁴, Pipr bowel syndrome: A pilot study Simpson², B U, Li¹

A multicenter registry study on percutaneous electrical nerve field stimulation for pediatric disorders of gut-brain interaction

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Ashish Chogle<sup>1</sup>
                        Khalil El-Chammas<sup>2</sup> | Neha Santucci<sup>2</sup> | Monica Grimm<sup>3</sup>
Lev Dorfman<sup>2</sup>
                      Kahleb Graham<sup>2</sup> | Daniel R. Kellv<sup>4</sup> | Jason E. Dranove<sup>4</sup>
                       Samuel Nurko<sup>5</sup> | Joseph Croffie<sup>6</sup>
Rachel Rosen<sup>5</sup>
Keshawadhana Balakrishnan<sup>7</sup> │ Eric H. Chiou<sup>7</sup> │ Liyun Zhang<sup>3</sup> │
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Neurostimulation for abdominal pain-related functional gastrointestinal disorders in adolescents: a randomised, double-blind, sham-controlled trial

Katja Kovacic, Keri Hainsworth, Manu Sood, Gisela Chelimsky, Rachel Unteutsch, Melodee Nugent, Pippa Simpson, Adrian Miranda

Impact of auricular percutaneous electrical nerve field Children with Cyclic Vomiting Syndrome stimulation on gut microbiome in adolescents with irritable

Geetanjali Bora, Samantha N. Atkinson, Amy Pan, Manu Sood, Nita Salzman, Katja Karrento 🔀

Impaired Vagal Efficiency Predicts Auricular Neurostimulation Response in Adolescent Functional Abdominal Pain Disorders

Katia Kovacic, MD1, Jacek Kolacz, PhD2-3, Gregory F. Lewis, PhD2-4 and Stephen W. Porges, PhD3-6

Percutaneous electrical nerve field stimulation for adolescents with irritable bowel syndrome: Cost-benefit and cost-minimization analysis

Eric Shah 🔀 Shanti Eswaran, Kimberly Harer, Allen Lee, Borko Nojkov, Prashant Singh, William D. Chey

Mınımal adverse effects profile following implantation of periauricular percutaneous electrical nerve field stimulators: a retrospective cohort study

Arthur Roberts 1, Alec Sithole 2, Marcos Sedghi 3, Charles A Walker 4, Theresa M Quinn 5

Emicacy of Auricular Neurostimulation in Adolescents With Irritable Bowel Syndrome in a Randomized, Double-Blind Trial

Amornluck Krasaelap,* Manu R. Sood, B U. K. Li,* Rachel Unteutsch,* Ke Yan, Melodee Nugent, Pippa Simpson, and Katja Kovacic

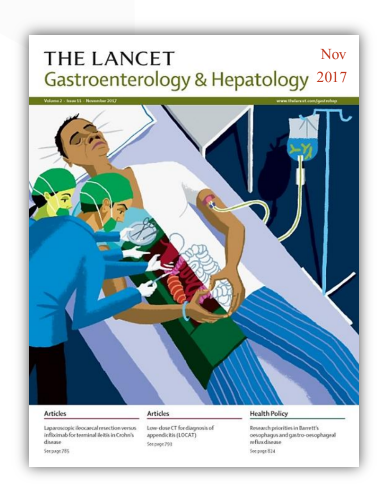
Impaired Vagal Efficiency Predicts Auricular Neurostimulation Response in Adolescent Functional Abdominal Pain Disorders

Katja Kovacic, MD1, Jacek Kolacz, PhD2-3, Gregory F. Lewis, PhD2-4 and Stephen W. Porges, PhD3-

Am J Gastroenterol 2020;115:1534-1538. https://doi.org/10.14309/ajg.00000000000000753

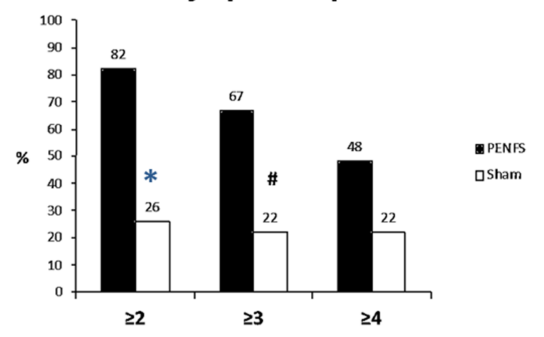
Growing Body of Clinical Evidence





Improvement of Global Symptoms in Patients with Irritable Bowel Syndrome

Global Symptom Improvement



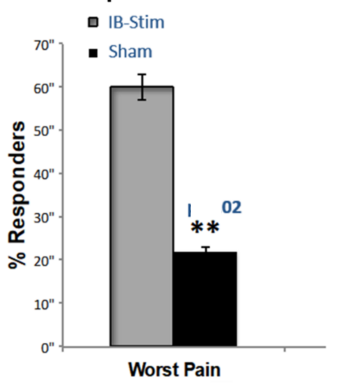
^{*}Percent of patients reporting improvement of global symptoms using Symptom Response Scale score ≥ 2 (*p ≤ 0.001), ≥ 3 (#p=0.002) and ≥ 4 (p=0.077)



FDA Benchmark for Clinically Meaningful Endpoint



≥30% Improvement in Pain



Treatment for abdominal pain-related functional gastrointestinal disorders in adolescents:

Number Needed to Treat (NNT):

The number of patients that need to be treated for one patient to get the targeted improvement (≥30% improvement).

IB-Stim NNT=3

IBS drugs in adults (lubiprostone, linaclotide, and rifaximin) NNT=6 to 14²

^{1.}Krasaelap A, et al. Efficacy of Auricular Neurostimulation in Adolescents With Irritable Bowel Syndrome in a Randomized, Double-Blind Trial. Clinical Gastroenterology & Hepatology. Clin Gastroenterol Hepatol. 2020;(9):1987-1994

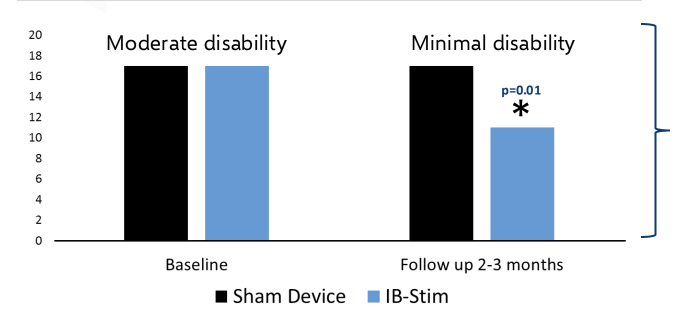




Functional Disability Scores at Long-Term Follow-Up



Improvement in Functional Disability in Patients with Irritable Bowel Syndrome



Improving functional disability (attending school and activities) is a marker of overall health and clinically meaningful beyond subjective pain measures

* Based on functional disability index (FDI) developed and validated tool to assess difficulties in daily functioning due to chronic pain.



Largest Pediatric Registry in Children with DGBI



A multicenter registry study on percutaneous electrical nerve field stimulation for pediatric disorders of gut-brain interaction

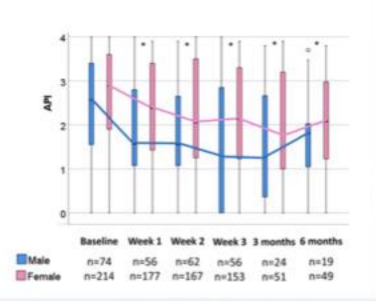
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Ashish Chogle<sup>1</sup> | Khalil El-Chammas<sup>2</sup> | Neha Santucci<sup>2</sup> | Monica Grimm<sup>3</sup> | Lev Dorfman<sup>2</sup> | Kahleb Graham<sup>2</sup> | Daniel R. Kelly<sup>4</sup> | Jason E. Dranove<sup>4</sup> | Rachel Rosen<sup>5</sup> | Samuel Nurko<sup>5</sup> | Joseph Croffie<sup>6</sup> | Keshawadhana Balakrishnan<sup>7</sup> | Eric H. Chiou<sup>7</sup> | Liyun Zhang<sup>3</sup> | Pippa Simpson<sup>3</sup> | Katja Karrento<sup>3</sup> |
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- Pediatric registry with "real world" clinical data
- 61% had failed ≥ 4 medication prior to treatment
- Sustained efficacy in abdominal pain up to 6-12 months after 4 weeks of IB-Stim treatment

A multicenter registry study on percutaneous electrical nerve field stimulation for pediatric disorders of gut-brain interaction. J Pediatr Gastroenterol Nutr. 2024

TABLE 1 Participating centers

Participating centers				
Center	Number of patients			
Cincinnati Children's Hospital	89			
Children's Hospital of Orange County	75			
Children's Wisconsin	65			
Atrium Health Levine Children's Hospital	31			
Boston Children's Hospital	18			
Riley Hospital for Children	11			
Texas Children's Hospital	3			





Percutaneous electrical nerve field stimulation compared to standard medical therapy in adolescents with functional abdominal pain disorders

Neha R. Santucci^{1,2*}, Rashmi Sahay³, Khalil I. El-Chammas^{1,2}, Kahleb Graham^{1,2}, Mikaela Wheatley^{1,2}, Madeleine Vandenbrink², Jennifer Hardy¹ and Lin Fei³

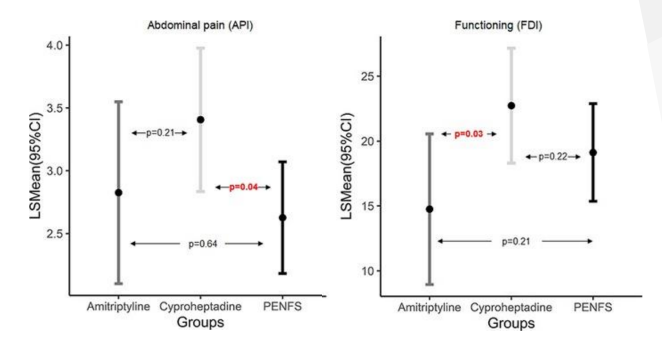
IB-Stim was equivalent or better than standard medications used for FAPDs

TABLE 2 Changes in measures in each group.

Treatment	Measure	Visit	LS means (LCL, UCL)	Diff LS means (LCL, UCL) ^a	<i>p</i> -value
PENFS	API	Baseline	2.776 (2.398, 3.153)		
		3 mFU	2.006 (1.512, 2.499)	-0.77 (-1.169, -0.371)	0.001
	NSS	Baseline	2.45 (2.039, 2.861)		
		3 mFU	1.738 (1.01, 2.466)	-0.712 (-1.456, 0.032)	0.059
	FDI	Baseline	20.244 (16.09, 24.399)		
		3 mFU	14.382 (8.215, 20.55)	-5.862 (-11.652, -0.073)	0.048
Cypro-heptadine	API	Baseline	3.555 (2.77, 4.34)		
		3 mFU	3.252 (2.456, 4.049)	-0.303 (-1.022, 0.416)	0.377
	NSS	Baseline	2.603 (2.026, 3.181)		
		3 mFU	2.054 (1.463, 2.645)	-0.550 (-1.259, 0.160)	0.117
	FDI	Baseline	23.785 (19.161, 28.408)		
		3 mFU	20.604 (15.161, 26.047)	-3.181 (-8.053, 1.691)	0.185
Amitriptyline	API	Baseline	3.113 (2.045, 4.182)		
		3 mFU	2.3 (1.186,3.413)	-0.814 (-1.553, -0.074)	0.034
	NSS	Baseline	2.007 (1.192, 2.822)		
		3 mFU	1.445 (0.579, 2.311)	-0.562 (-1.262, 0.138)	0.101
	FDI	Baseline	15.944 (8.352, 23.537)		
		3 mFU	11.709 (2.597, 20.82)	-4.236 (-12.195, 3.723)	0.259

Examined using Chi square test.

PENFS, percutaneous electrical nerve field stimulation; API, abdominal pain index; NSS, nausea severity scale; FDI, functional disability inventory; LS, least square; LCL, lower control limit; UCL, upper control limit.



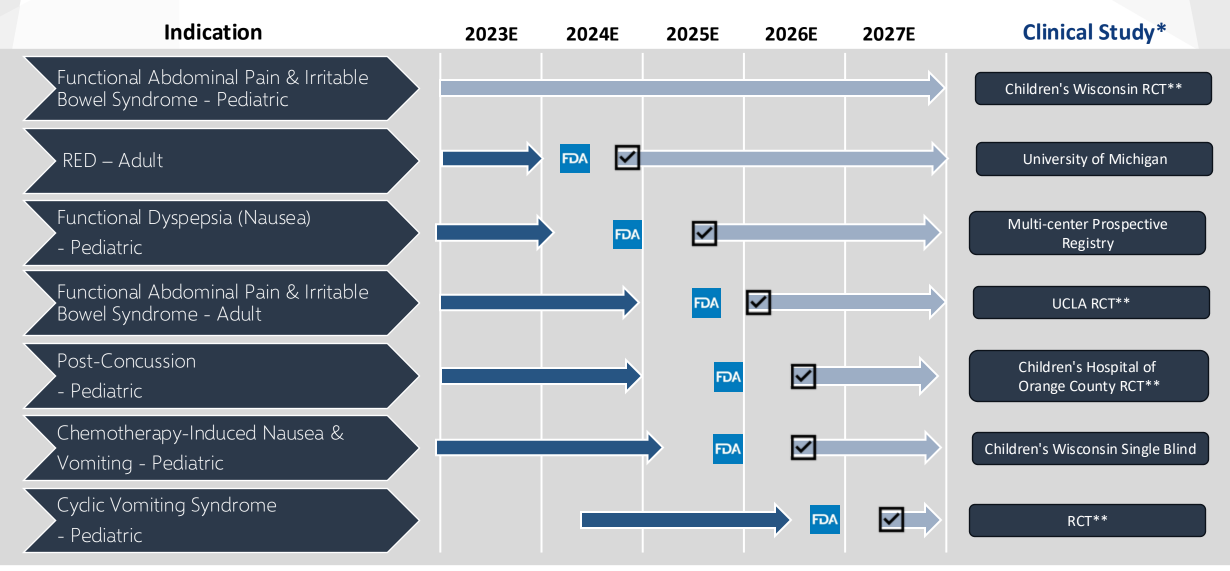


^{*}negative values indicate reduction in outcome scores from baseline to 3month Follow Up visit.

IB-Stim[™] vs. Drugs Competitive Landscape

		Antidepressants		Adult Use (Peripherally Acting at the Gut Level)			Gut Level)
	IB-Stim™	Amitriptyline	Citalopram	Amitiza	Linzess	Trulance	Viberzi
FDA Approved for IBS in Children and Adolescents	✓						
Improves Functional Disability	✓						
Targets Brain-Gut Axis	✓	✓	✓				
Better Than Placebo for Pain in IBS	✓			✓	✓	✓	✓
Improves Pain Catastrophizing	✓						
Improves Global and Somatic Symptoms	✓						
Most Serious Potential Side Effects	Localized Skin Irritation	Suicidal Ideation, Dementia (long term use)	Suicidal Ideation, Dementia (long term use)	Abdominal Pain, Allergic Reaction	Diarrhea, Abdominal Pain	Diarrhea, Serious Allergic Reaction	Pancreatitis, Serious Allergic Reaction, Intestinal Obstruction
Easily Accessible	✓	✓	✓	✓	✓	✓	✓

FDA Pipeline - Indications and Timelines













Expanding Portfolio of Next Generation Devices for Disorders of Gut-Brain Interaction



RECTAL EXPULSION DEVICE [RED]

- Developed at the University of Michigan enabling comprehensive constipation care for every adult gastroenterology practice
- RED is a self-inflating balloon expulsion test that allows for point-of-care testing to effectively identify patients with an evacuation disorder
- FDA 510(k) submission on track for June 2024 with expected clearance in Q4 2024



RESPONSIVE DESIGN





MARKET

- ~\$2B market opportunity
- Current balloon expulsion testing requires a separate visit to a GI physiology laboratory
- Anorectal manometry is too expensive to be practical



Self-inflating



- Enables point-of-care testing (In-office use)
- Provides immediately actionable binary test results



REIMBURSEMENT

- Current CAT I CPT Code 91120
- Medicare reimbursement: \$519.15



IB-Stim Reimbursement Market Access Plan Established

Lives



Strong Data = Strong Policy Coverage & Reimbursement = Strong Revenue Growth

Key Publications		Academic Society Support		Successful Cov	verage Policies		CPT Billing Code
• Versus Placebo	• NA	ASPGHAN Written Support		• CareFirst BCBS	BCBS Kansas City	• CAT III (E	ffective July 2022)
• Long-term Data x 2	• A/	AP Written Support		• BSBS Nebraska	• Highmark BCBS	• CAT I App	plication (June 2024)
• Comparison vs. Soc Rx				• Quartz Wisconsin	• FL Blue	• CAT I Co	de (Effective January 2026
Multi-center Registry				BCBS South Carolin	a • BCBS VT		
o ~300 patients				BCBS Massachusett	_s • Geisenger		
	_				BCBS of North Dako	nt a	
 Positive Health Economic 	c Data			 CS HealthVine 	DCD3 OF NOTHER DAKE	λία	
Positive Health Economic 2023E	c Data	2024E		• CS HealthVine	2020		2027E
2023E	c Data		A Indicatio RED	2025E	2020		2027E
	c Data	700+ Published Patients 16		2025E	FDA FDA Indication: Adult FAP/IBS	FDA Indication: Post-Concussion Syndrome	2027E FDA FDA Indication Cyclic Vomiting Syndrome
2023E	c Data	700+ Published Patients 16		2025E	FDA FDA Indication: Adult FAP/IBS	FDA Indication: Post-Concussion Syndrome Post-Concussion Syndrome	FDA FDA Indication Cyclic Vomiting

IB-Stim Go-to-Market Strategy

Strong Data = Strong Policy Coverage & Reimbursement = Strong Revenue Growth

2024 Policy Coverage

Total Plans	Total # Lives covered
12 plans	~24M

Insurance Plans	# Lives covered
CareFirst BCBS	3.5M
BCBS of MA	3M
BCBS of SC	770k
BCBS of Nebraska	340k
Quartz Wisconsin	335k
CareSource HealthVine	120k
BCBS Kansas City	1M
Highmark BCBS	7M
Florida Blue	6M
BCBS VT	200K
Geisenger	600K
BCBS North Dakota	310K
* CDT Code Effective July 1st 2022	

Commercialization Strategy



On strength of clinical evidence, targeting guideline changes that support IB-Stim™ as standard of care

Launched internal Prior Authorization Team

Increasing D2C Marketing in States with Policy Coverage

Hiring W-2 Reps in States with Policy Coverage



Reimbursement Strategy

Technology Specific CPT coding

recritiology specific of 1 county				
CPT code*	CAT III code (0720T)			
List Price	• \$1,195			

Engaging with AAP and NASPGHAN to apply for CAT I CPT code in 2024

Customers



~33k

U.S. Pediatricians

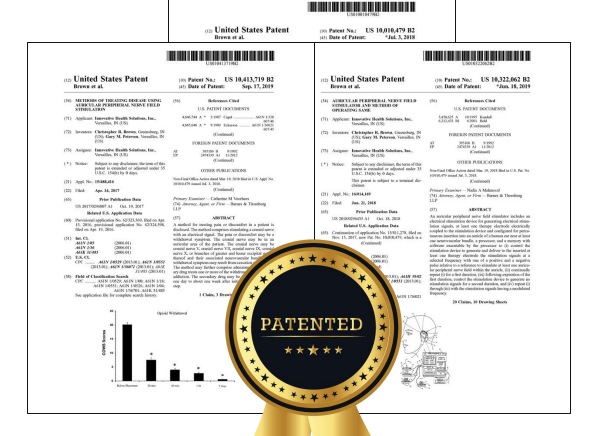
~10k

U.S. Adult Gastroenterologists



NeurAxis IP Portfolio

- 11 issued and 9 pending patents
 - Device
 - Method
- U.S. IP runs through 2039 as of now
- International IP in process
- Freedom to operate completed



Collaborative Contract Manufacturing Partner



In-House Capabilities

- Office and factory in Indiana
- 69,000 square foot facility
 - Offices, factory, environmentally controlled room, warehouse, parts processing, assembly, quality control
- Medical device focused manufacturing established in 1990

Manufacturing Capacity

- Controlled, repeatable, monitored production process
- Kit production capacity sufficient for all NeurAxis projected needs
- New dedicated room built in 2022 for NeurAxis equipment and production
 - All NeurAxis materials now maintained in the room
 - Environmentally controlled build room

Quality Management System

- ISO 13485:2016 Certified
- FDA registered
- ITAR Registered











Medical Advisory Board





Dr. Samuel Nurko





Dr. Carlo Di Lorenzo





Dr. Rachel Rosen





Dr. Kahlil El-Chammas





Dr. Miranda van Tilburg





Dr. Leonel Rodriguez





Board of Directors





Beth Keyser Board Member



Mitch Watkins Board Member



Kirstin Ferge Board Member



Brian Carrico Chief Executive Officer, **Board Member**



Dr. Chris Brown Director Of Innovation, Founder, Board Member

Collective Experience

























Management Team





Brian Carrico
Chief Executive Officer,
Board Member







Dr. Adrian Miranda *Chief Medical Officer*







Timothy Henrichs
Chief Financial Officer









Dr. Tom CarricoChief Regulatory Officer





Dr. Chris BrownDirector of Innovation,
Founder, Board Member



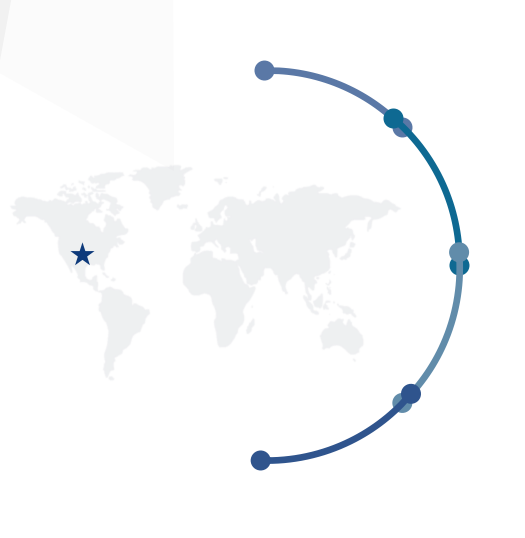






Key Investment Highlights

Strong Data = Strong Policy Coverage & Reimbursement = Strong Revenue Growth





Large Global Market with Significant Unmet Need

- \$23B+ TAM³ for target pipeline indications
- \$9B+ TAM³ for target pediatric indications (near-to-mid term)
- \$14B+ TAM³ for target adult indications (mid term)
- Large unmet clinical need: high refractory, off label pharmacological treatments with adverse side effects



Unique, Innovative Product Supported by Clinical Evidence

- Novel treatment targeting the brain-gut-axis
- Differentiated PENFS technology
- 700+ published patients⁴
- Easy-to-learn and efficient procedure



Clear Commercial Pathway

- FDA De Novo clearance
- Technology-specific CPT billing code
- Major Insurance Payer Coverage initiated
- Strong IP on Device and Method



Seasoned Management and Board

- Experienced management team and Board of Directors
- Operations and infrastructure built to scale
- Path to profitability

