

A Commercial-Stage Bioelectronic Medicine & Wellness Company

Nasdaq: ECOR

Corporate Presentation

October 2024

Forward Looking Statement

In addition to historical information, this presentation may contain forward-looking statements with respect to our business, capital resources, strategy and growth reflecting the current beliefs and expectations of management made pursuant to the safe harbor provisions of the Private Securities Litigation Reform Act of 1995. Forward-looking statements are subject to a number of risks, uncertainties and assumptions, and you should not rely upon forward-looking statements as predictions of future events. All forward-looking statements may be based upon current estimates and expectations about future events and financial and other trends. There is no guarantee that future results, performance or events reflected in the forward-looking statements will be achieved or occur. No person assumes responsibility for the accuracy and completeness of the forward-looking statements, and, except as required by law, we assume no obligation to update these forward-looking statements publicly, or to update the reasons why actual results could differ materially from those or our situation may change in the future.

Forward-looking statements include all statements that are not historical facts and, in some cases, can be identified by terms such as "anticipates," "believes," "could," "seeks," "estimates," "intends," "may," "plans," "potential," "predicts," "projects," "should," "will," "would" or similar expressions and the negatives of those terms. Forward-looking statements involve known and unknown risks, uncertainties and other factors that may cause our actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements. Forward-looking statements represent our management's beliefs and assumptions only as of the date they are made and are only predictions that may be inaccurate. You should read the Risk Factors set forth in our reports filed from time to time with the Securities and Exchange Commission, which factors may cause our actual future results to be materially different from what we expect.







gammaCore



1st FDA-cleared non-invasive vagus nerve stimulator

- Fast acting, comfortable, easy to use hand-held option
- FDA Cleared for the prevention and treatment of primary headache in adults and adolescents
- No drug-drug interactions or drug-like side effects
- Can be used as a stand-alone therapy or alongside existing treatments
- Can use multiple times per day or month
- Cost dominance in the first-year when gammaCore therapy is used in conjunction with standard of care as supported by UK NICE guidance

Investment Summary

Platform Therapy	FDA cleared; proprietary, non-invasive vagus nerve stimulator (nVNS) positioned to unlock the broad potential of bioelectronic medicine	
Consistent Topline Growth	Approximately \$16M revenue in 2023; 5-year revenue trailing twelve month (TTM) compounded annual growth rate (CAGR%) of 69% Gross margin consistently above 80%	
Diverse Market Opportunity	Primary Headache estimated to affect more than 39 million ¹ adults in the U.S. Expanding into Wellness and Human Performance to take advantage of the \$5 trillion Digital Health and Wellness market ²	
Attractive Business Models	Variety of products, price points, and business models for different end users	
Strong IP Portfolio	Patent coverage extends beyond 2037 1. American Migraine Foundation. 2. Digital Health & Wellness Market Size Trends Report, Forecast.	



Benefits of nVNS (Non-invasive Vagus Nerve Stimulation)

The vagus nerve affects multiple organs and systems

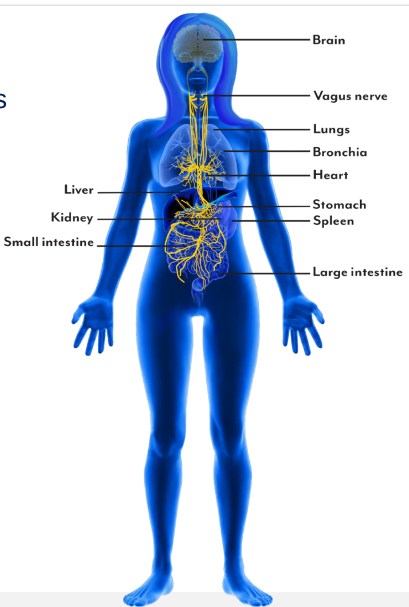
Activates multiple mechanisms of action

Evidence supports a variety of current and future treatment targets



Personal use, pocket size, portable, easy to use, products





CNS: Reduces nociception, reduces cortical spreading depression, and other brain modulation effects; increases glymphatic flow

Cardiac: Reduces heart rate and blood pressure

Pulmonary: Increases bronchodilation

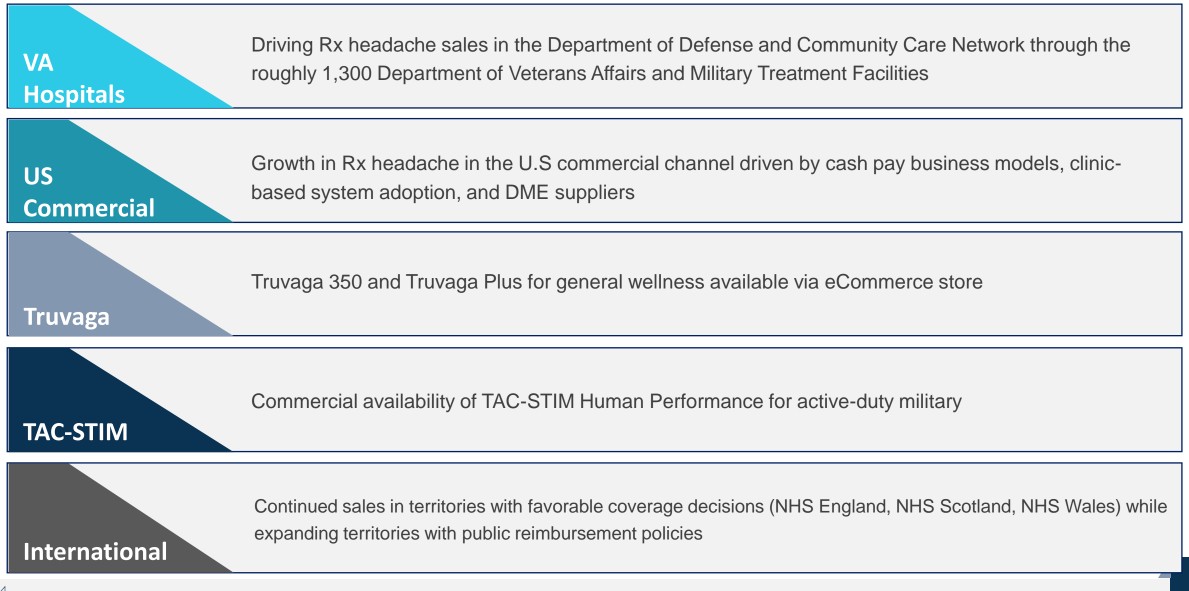
Hepatic: Regulation of gluconeogenesis

Gastrointestinal: Increases GI motility and secretions; satiation

Splenic: Detection and regulation of systemic inflammation

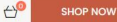


Active Channels With Revenue Growth Opportunities



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electro**Core**



LEARN MORE

Meet the Truvaga Vagus Nerve Stimulators



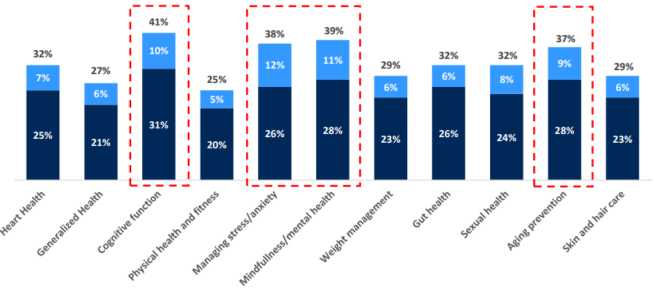
Truvaga[™] is a general wellness product and as such has not been evaluated by the US Food and Drug Administration. Truvaga products are not intended to diagnose, treat, cure, or prevent any disease or condition.





Cognitive Function and Stress Remain an Unmet Need for Consumer

Global stress market is estimated at \$20.6 billion with the U.S. stress market growing at a 5.3% CAGR(%)¹



Some products/services available but wish there were more

Not nearly enough products/services available

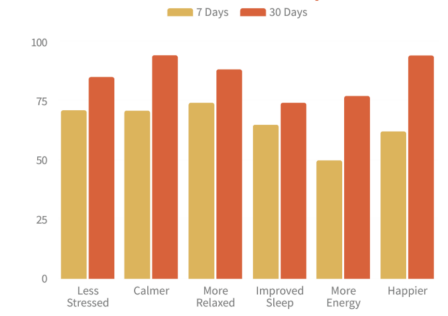
Source: McKinsey Future of Wellness Survey, August 2023

Source: Truvaga Consumer Study

*Based on 30-day consumer in-home use test performed by The Benchmarking Company, an independent third-party research firm. The study consisted of 34 participants using Truvaga morning and night for 30 days.



1. <u>https://www.grandviewresearch.com/industry-analysis/us-workplace-stress-management-market-report</u>. CAGR(%) is 2023 – 2030.



The Benefits of Daily Use*

electro**Core**



TAC-STIM

LEARN MORE

TAC-STIM has been developed with support from the 711th Human Performance Wing Performance Optimization Branch of the United States Air Force to provide:

- Accelerated Training
- Sustained Attention
- Reduced Fatigue
- Improved Mood

FAC-STIM

E IN THE USA

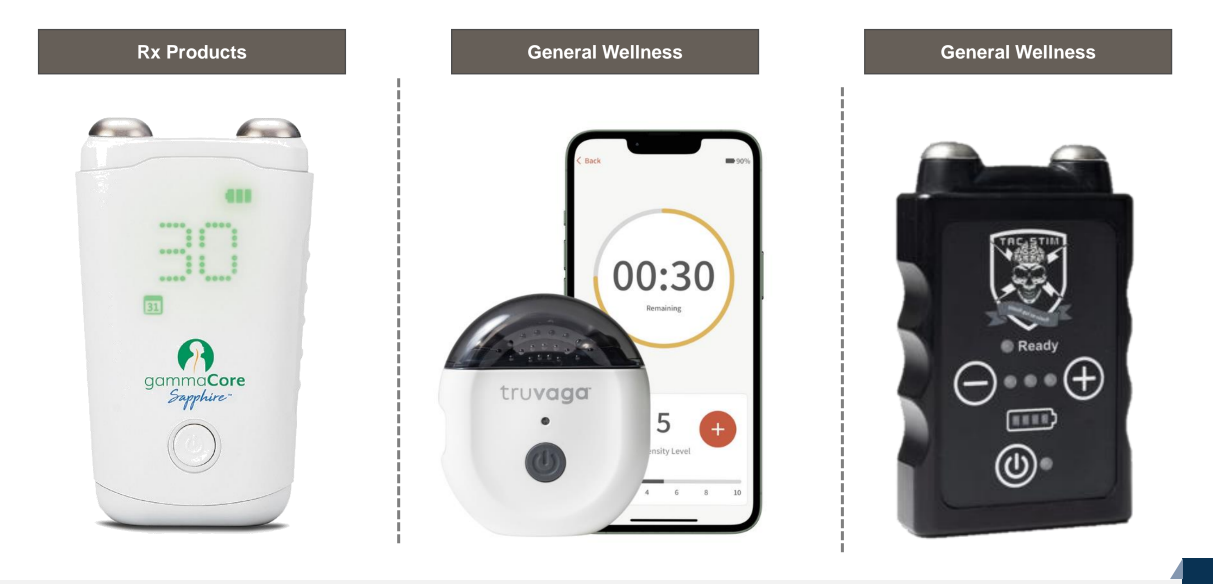
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9

Ready

nVNS – A Platform Technology









- Post-traumatic stress disorder^{2,3}
- Opioid Use Disorder²
- Traumatic Brain Injury²
- Parkinson's Disease²
 - Acute Stroke²



gammaCore is the only FDA-cleared non-invasive VNS therapy in primary headache

¹ Cleared indications, ² Investigator Initiated Trials ongoing, ³Breakthrough Designation

gammaCore (nVNS) currently is FDA-cleared for prevention of migraine and cluster headache, and acute treatment of migraine and episodic cluster headache, as well as the acute and preventive treatment of migraine in adolescents between 12 and 17 years of age



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Broad Intellectual Property Portfolio

electroCore owns all intellectual property on which the technology relies

Expansive pioneering IP coverage of non-invasive, transdermal neuro-stimulation in the neck

We have patent coverage extending beyond 2037:

- High-frequency burst signals capable of passing comfortably through the skin
- Low-pass signal filtration that reduces signal harmonics that cause pain
- Growing digital health portfolio

>200

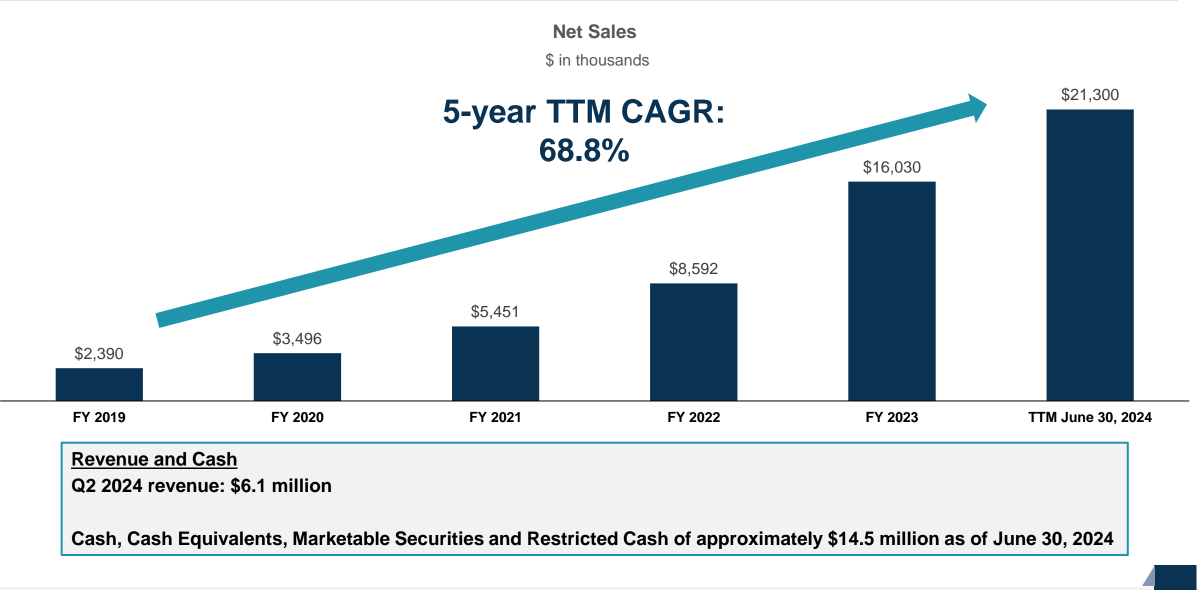
PATENTS AND PATENT APPLICATONS

~140 issued U.S. patents

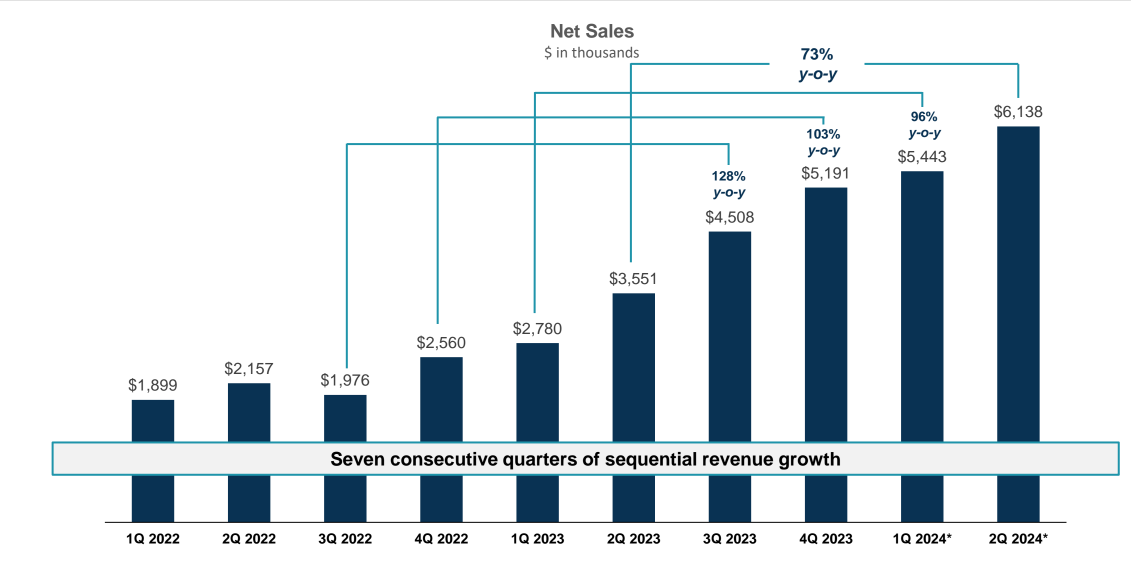
- ~30 U.S. patent applications
- ~35 International patents and applications



Summary Financials and Guidance



Summary Revenue by Quarter

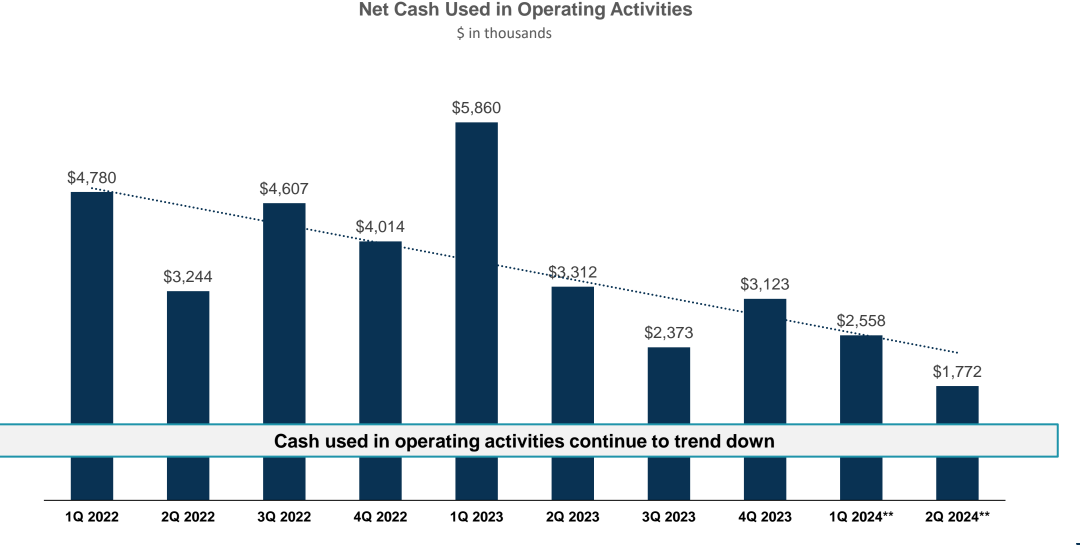


*Unaudited Actuals



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Summary Net Cash Used in Operating Activities



**Unaudited Actuals



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Capitalization Table

Fully diluted as of June 30, 2024 (in thousands)

Common Shares	6,447	
Pre-Funded Warrants	1,608	
Warrants	1,640	Average Exercise price = \$5.37
Options	498	Average Exercise price = \$36.98
Restricted and Deferred Stock Units	422	RSUs/DSUs which vest through January 2027
Total	10,615	



Experienced Management Team





Investment Summary

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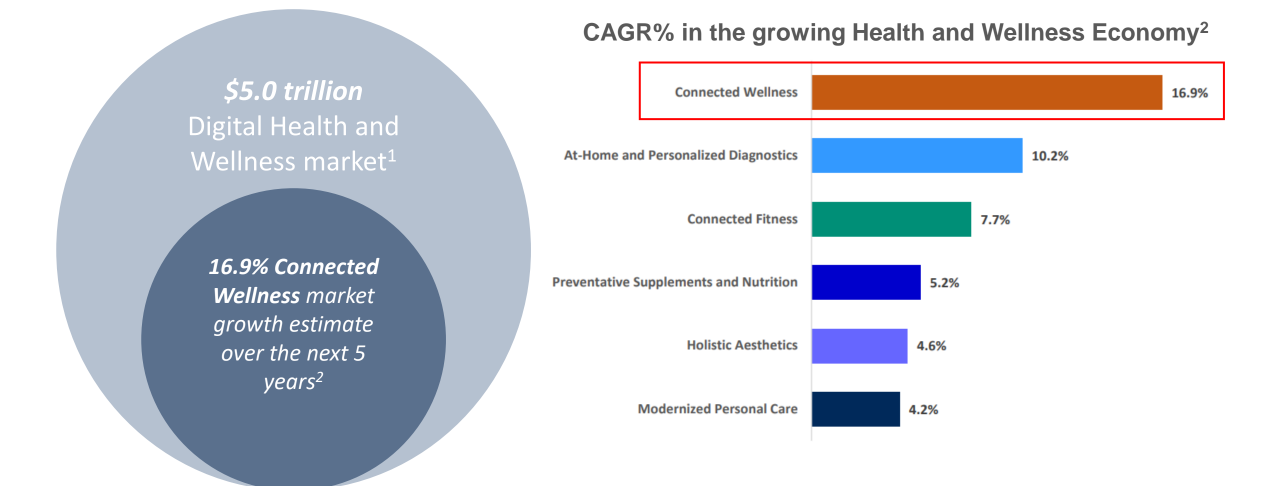


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Leveraging nVNS in the \$5 trillion Digital Health and Wellness Market



1. Digital Health & Wellness Market Size Trends Report, Forecast.

2. Source: Canaccord Genuity Industry Update: Health, Wellness and Lifestyle. Figure 4. January 4, 2024.



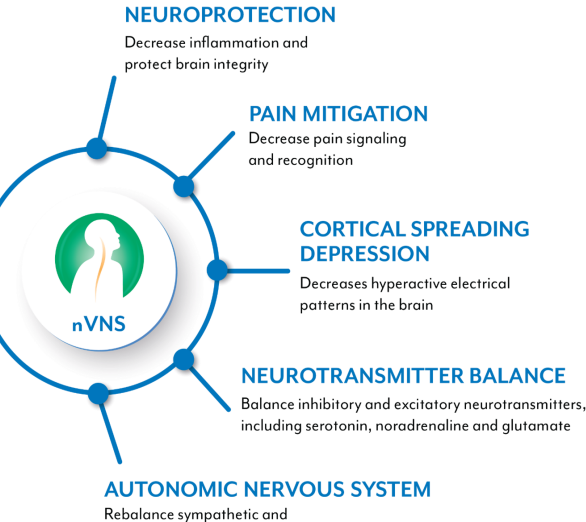


Mechanism of Action

nVNS: An Integrative Mechanism of Action

There are multiple known mechanism of vagus nerve stimulation:





parasympathetic input



Global Approvals

FDA-Cleared Indications for the US

- The preventive treatment of migraine headache in adolescent (age 12 and older) and adult patients.
- The acute treatment of pain associated with migraine headache in adolescent (age 12 and older) and adult patients.
- Adjunctive use for the preventive treatment of cluster headache in adult patients.
- The acute treatment of pain associated with episodic cluster headache in adult patients.
- Treatment of hemicrania continua in adults.
- Treatment of paroxysmal hemicrania in adults.

US FDA guidance document titled: "General Wellness: Policy for Low-Risk Devices, Guidance for Industry and Food and Drug Administration Staff"

Meets the following two factors, and,

- Is intended for only general wellness use
- Presents a low risk to the safety of users and other persons

It's intended uses involve claims about sustaining or offering general improvement to functions associated with a general state of health that do not make any reference to diseases or conditions. Examples of these include, but are not limited to "claims" related to:

- relaxation or stress management (including claims to promote relaxation or manage stress)
- mental acuity (including claims to improve mental acuity, instruction following, concentration, problem-solving, multitasking, resource management, decision-making, logic, pattern recognition, or eye-hand coordination, as well as enhancing learning capacity)

CE Marks for the EU/EFTA/EEA and UK

- Acute and/or prophylactic treatment of primary headache (migraine, cluster headache, and hemicrania continua) and medication overuse headache in adults.
- Treatment or prevention of symptoms of reactive airway disease, including asthma, bronchoconstriction, exercise-induced bronchospasm, and COPD.
- Adjunctive therapy for adults to reduce the symptoms of certain anxiety and depression conditions (including panic disorder, posttraumatic stress disorder, obsessive-compulsive disorder, and major depressive disorder).
- Adjunctive therapy in the prevention of partial onset and generalized seizures associated with epilepsy in adults.
- Adjunctive therapy for adults to reduce the symptoms of gastric motility disorders and irritable bowel syndrome (including nausea, vomiting, bloating/distention, early satiety, and abdominal pain).

Health Canada License for Canada

Acute and/or prophylactic treatment of migraine in adolescents and adults and cluster headache in adults.

ARTG Certificate for Australia

Acute and/or prophylactic treatment of migraine, cluster headache, and hemicrania continua in adults.





Rx gammaCore Important Safety Information

gammaCore SapphireTM (non-invasive vagus nerve stimulator) is intended to provide non-invasive vagus nerve stimulation (nVNS) on the side of the neck. gammaCore is indicated for:

- The preventive treatment of migraine headache in adolescent (age 12 and older) and adult patients.
- The acute treatment of pain associated with migraine headache in adolescent (age 12 and older) and adult patients.
- Adjunctive use for the preventive treatment of cluster headache in adult patients.
- The acute treatment of pain associated with episodic cluster headache in adult patients.
- Treatment of hemicrania continua in adults.
- Treatment of paroxysmal hemicrania in adults.

The effectiveness of gammaCore (nVNS) has not been established in the acute treatment of chronic cluster headache.

The long-term effects of the chronic use of the device have not been established.

gammaCore contraindications include but are not limited to:

- · Patients that have an active implantable medical device, such as a pacemaker, hearing aid implant, or any implanted electronic device
- Patients that have a metallic device such as a stent, bone plate, or bone screw implanted in or near the neck
- Patients that are using another device at the same time (e.g., TENS unit, muscle stimulator) or any portable electronic device (e.g., cell phone).

Safety and efficacy of gammaCore have not been evaluated in the following patients, and therefore gammaCore is NOT indicated for:

- · Adolescent patients with congenital cardiac issues
- Patients diagnosed with narrowing of the arteries (carotid atherosclerosis)
- Patients who have undergone surgery for resection of the vagus nerve in the neck (cervical vagotomy)
- Pediatric patients (less than 12 years)
- Pregnant women
- · Patients with clinically significant hypertension, hypotension, bradycardia or tachycardia

The long-term safety and effectiveness of the gammaCore device has not been demonstrated in adolescents 12-17 years of age. Due to hormonal and cognitive development changes in adolescents, this population should be closely monitored while using the device. The use of the device in this population is based on extrapolated data from a clinical study in adults.

- You must read the gammaCore Instructions for Use before using gammaCore
- · Only use gammaCore as described in these Instructions for Use or as otherwise directed by your Healthcare Provider
- Only use an electroCore-approved electrode gel with gammaCore.

In the US, the FDA has not cleared gammaCore for the treatment of pneumonia and/or respiratory disorders, such as acute respiratory stress disorder associated with COVID-19. Please refer to the gammaCore Instructions for Use for all of the important warnings and precautions before using or prescribing this product: <u>www.gammacore.com</u> Please also see the instructions for Use for gammaCore CV for all the important warnings and precautions specific to gammaCore CV and its use pursuant to the Emergency Use Authorization (EUA): <u>https://www.fda.gov/media/139970/download</u>



Truvaga Warnings and Precautions

Warnings

Do not use Truvaga if:

- You have an active implantable medical device, such as a pacemaker, hearing aid implant, or any implanted electronic product.
- You are using another product at the same time (e.g., TENS Unit, muscle stimulator).
- You are driving, operating machinery, or during any activity that may put patient at risk of injury.
- You are in the presence of strong electromagnetic fields, such as MRI scanners.
- You are in an explosive atmosphere or in the presence of flammable gas mixtures.
- You have an open wound, rash, infection, swelling, cut, sore, drug patch, or surgical scar(s) on the neck at the application site.
- You have wet skin, are in the water, or just stepped out of the water (e.g., shower, bath, pool).

It is not recommended to use Truvaga:

- If you have had surgery to cut the vagus nerve at the neck as it may not be effective.
- If you are younger than 12.
- If you are pregnant or breastfeeding.
- More than 24 times a day.

Precautions

- Truvaga should be used only as described in the User Manual.
- Only use a Truvaga-supplied conductive gel.
- Do not apply Truvaga across or through the head, directly on the eyes, covering the mouth, on the chest, on the upper back, or over the heart.
- Do not use Truvaga if there are signs of damage or defects.
- Do not use if the light is flashing green and "Err" is displayed on the screen when the product is turned on.
- Do not submerge Truvaga in water; it is not water resistant.
- Store in a safe location out of reach of children.

Users with sensitive skin may experience application site discomfort, irritation and/or redness. If you experience light-headedness, dizziness, chest pain, excessive skin irritation, local pain, face/ head/neck area (including toothache), excessive muscle twitching, tingling, contractions, or other adverse reactions, DISCONTINUE USE. These reactions typically resolve after the session is complete; however, if it persists after the session, consult your physician.

Truvaga[™] are <u>general wellness product</u>s and as such have not been evaluated by the US Food and Drug Administration. Truvaga products are not intended to diagnose, treat, cure, or prevent any disease or condition.



TAC-STIM Warnings and Precautions

Warnings

Do not use Truvaga or TAC-STIM if:

- You have an active implantable medical device, such as a pacemaker, hearing aid implant, or any implanted electronic product.
- You are in an explosive atmosphere or in the presence of flammable gas mixtures.
- You have an open wound at the application site.

Precautions

- TAC-STIM should be used only as described in the User Guide.
- Do not use TAC-STIM if there are signs of damage or defects.
- Do not use if an error code is displayed on the screen when the product is turned on.
- Do not submerge TAC-STIM in water; it is not water resistant.

NOTE: There is no data on the use of TAC-STIM if you are pregnant or breastfeeding.

TAC-STIMTM, non-invasive Vagus Nerve Stimulator (nVNS) is a Low-Risk General Wellness Product. The product is not regulated by the US FDA and is not intended to treat or diagnose any medical condition or disease.

