

A large, stylized graphic of a human brain in profile, facing right. The brain is composed of green and white geometric shapes, including triangles and circles, some of which resemble gears. The background is a light green gradient with scattered green and white dots and circles of varying sizes.

Enabling the Nervous System to Repair Itself

CORPORATE PRESENTATION

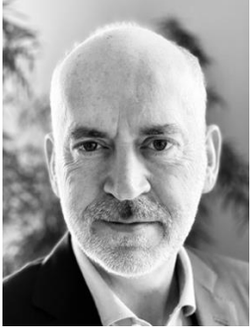
May 12, 2022

Financial Disclosure Statement

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NervGen's Clinical Trials are Led by Experienced Drug Developers



Paul Brennan
President & CEO

- 30+ years of biotech and pharma commercial and development experience
- Participated in the commercial or regulatory development of >10 products now EMA or FDA approved including budesonide (Pulmicort), esomeprazole (Nexium), budesonide/formoterol (Symbicort) and plerixafor (Mozobil)
- \$3+ billion in M&A, licensing and corporate restructuring transactions

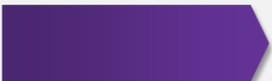


Dr. Dan Mikol, MD, PhD
Chief Medical Officer

- 25+ years pharma experience and as practicing neurologist conducting clinical research
- Joined NervGen from Amgen where he was Executive Director and Global Therapeutic Development Head, Neurology and Nephrology
- Participated in development and/or commercialization of natalizumab (Tysabri), fingolomod (Gilenya), cladribine (Mavenclad), interferon- β -1a and erenumab (Aimovig)

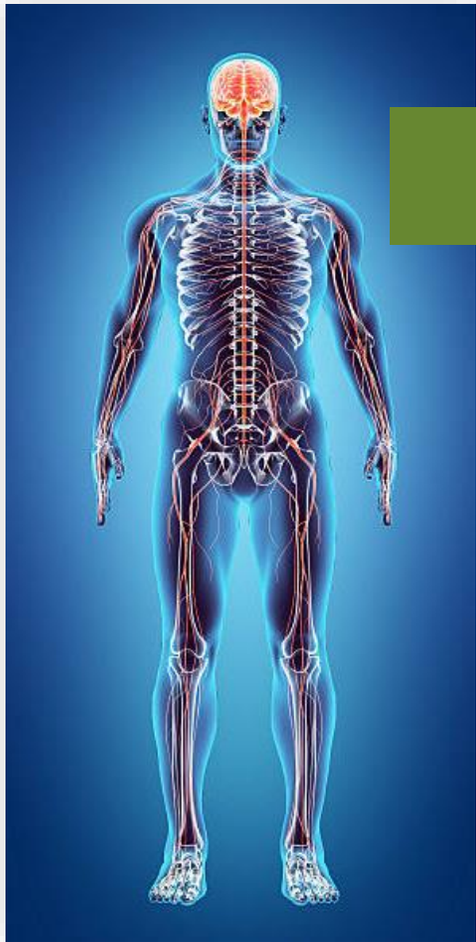


NVG-291: A Pipeline in a Product

INDICATION	STAGE OF DEVELOPMENT			ESTIMATED COST	MARKET OPPORTUNITY
	Phase 1	Phase 1b/2 Initiation	Phase 1b/2 Readout		
Alzheimer's Disease		Q4 2022	2024	\$20 M	<ul style="list-style-type: none"> • ~6,000,000 patients in the US • US Market potential of over \$300 billion • Substantial pharma deal dynamics
Spinal Cord Injury		Q4 2022	2023	\$10 M	<ul style="list-style-type: none"> • ~18,000 new patients per year in the US • ~300,000 chronic patients • Lifetime costs range from \$1 to >\$5 million
Multiple Sclerosis		Q1 2023	2024	\$20 M	<ul style="list-style-type: none"> • ~900,000 patients in the US • US Market potential of over \$30 billion • Currently there are multiple blockbusters

Proof of concept readouts for all three indications expected in 18-24 months

Revolutionizing the Treatment of Nervous System Damage



Everyone **KNOWS**...

The nervous system is a **complex system** that controls thought, movement, senses, etc.

Everyone **BELIEVES**...

The nervous system **cannot** repair itself



NVG-291 – First-in-Class *Neuroreparative* Drug

Representative of Placebo Group

(Back Legs and Tail Dragging)



Representative of NVG-291 Group

(Back Legs and Tail Active)



Remarkable and robust repair across multiple models

NervGen's Technology Was Invented by Dr. Jerry Silver

Known in the Spinal Cord Injury Field as the "Oracle"



Jerry Silver, PhD

Professor and Researcher,



Adjunct Professor,



Dr. Silver's Spinal Cord Research

- Discovered why the nervous system does not repair itself
- Identified the surprising molecules responsible

Dr. Silver Has Received Numerous Prestigious Awards Including

- Ameritec Prize
- Christopher Reeve-Joan Irvine Research Medal
- Jacob Javits Neuroscience Investigator Award

Dr. Silver's research revolutionized the understanding of the nervous system

STRONG IP PORTFOLIO

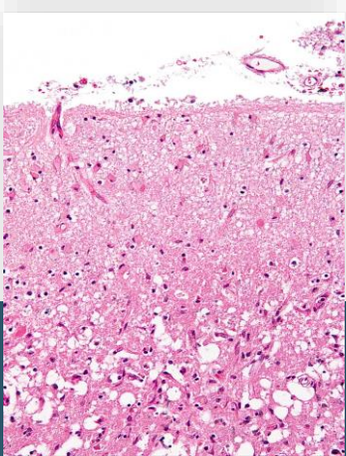
NervGen licensed the technology from Case Western and **owns global rights for all indications**

Intellectual property protection on NVG-291 until 2037

The Evolution of Our Proprietary Science

Pre 1990

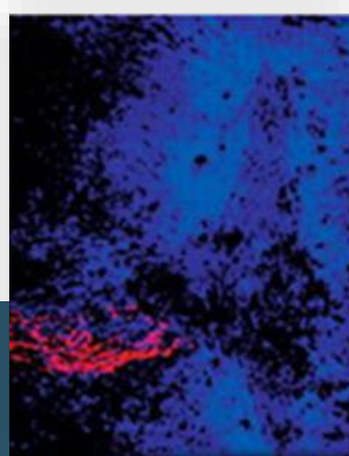
It was demonstrated that **glial scars** form at the site of injury to the nervous system and that scars in the brain cause neurons to be dysfunctional. Scars were later identified as the primary impediment of recovery



Micrograph of a glial scar

1990s

Dr. Silver identified a class of molecules called **CSPGs**, present in scars in the brain and spinal cord, that stop the body's natural repair mechanisms



Spinal cord nerve (red) trapped in the scar by CSPGs (blue)¹

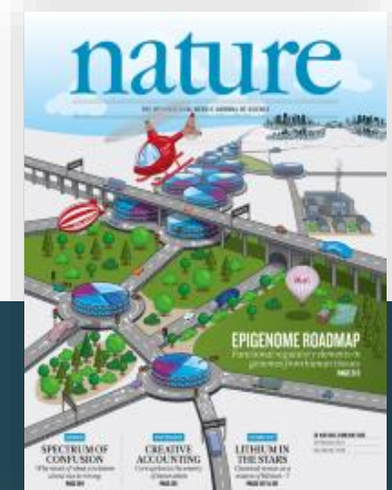
2009

Dr. Silver and collaborators from Harvard co-discovered that CSPGs bind with a receptor (**PTP σ**) present in the brain and spinal cord and that this interaction stops cells from repairing damage

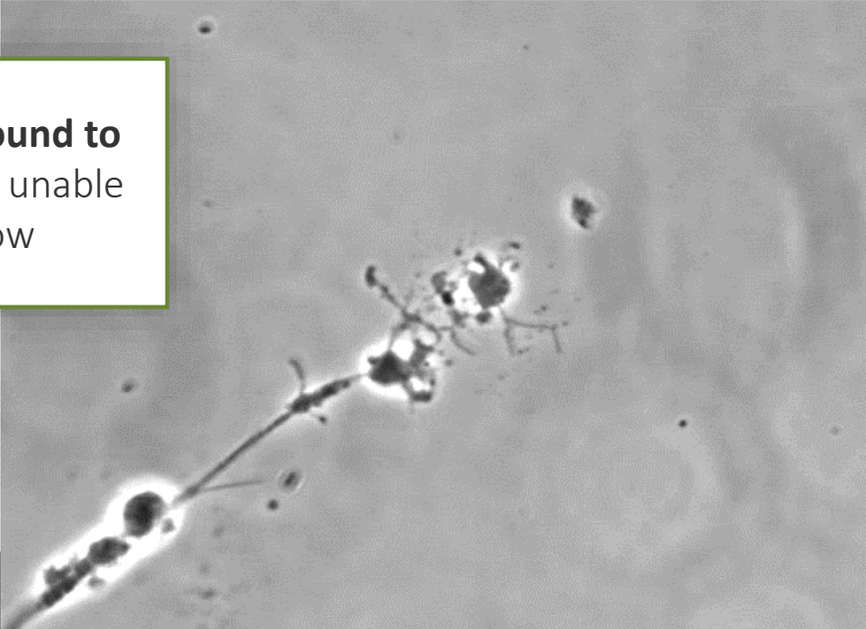


2015

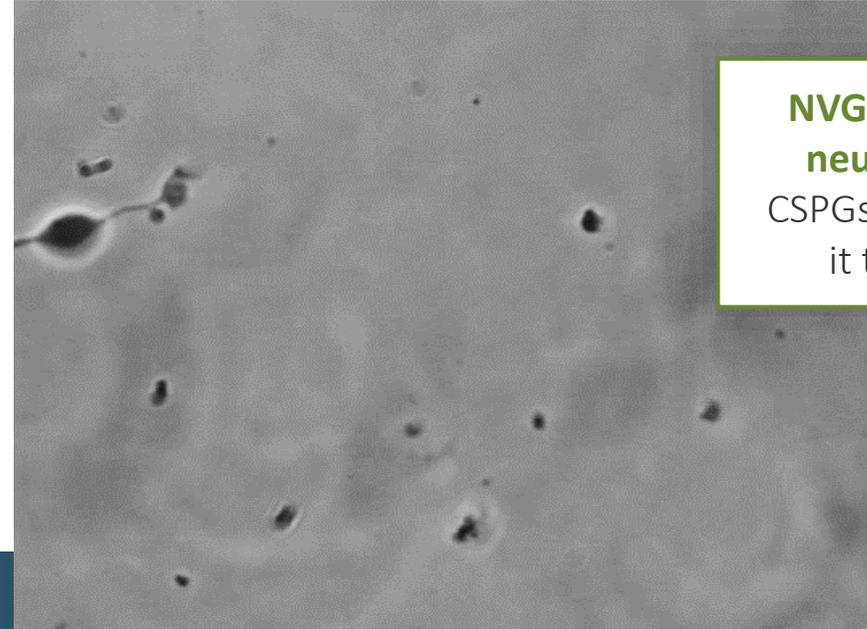
Dr. Silver's team then identified **NVG-291**, a drug that targets the interaction between CSPGs and PTP σ and allows the nervous system to repair damage



NVG-291 Allows Neurons to Grow in the Scar



Neuron bound to CSPGs and unable to grow



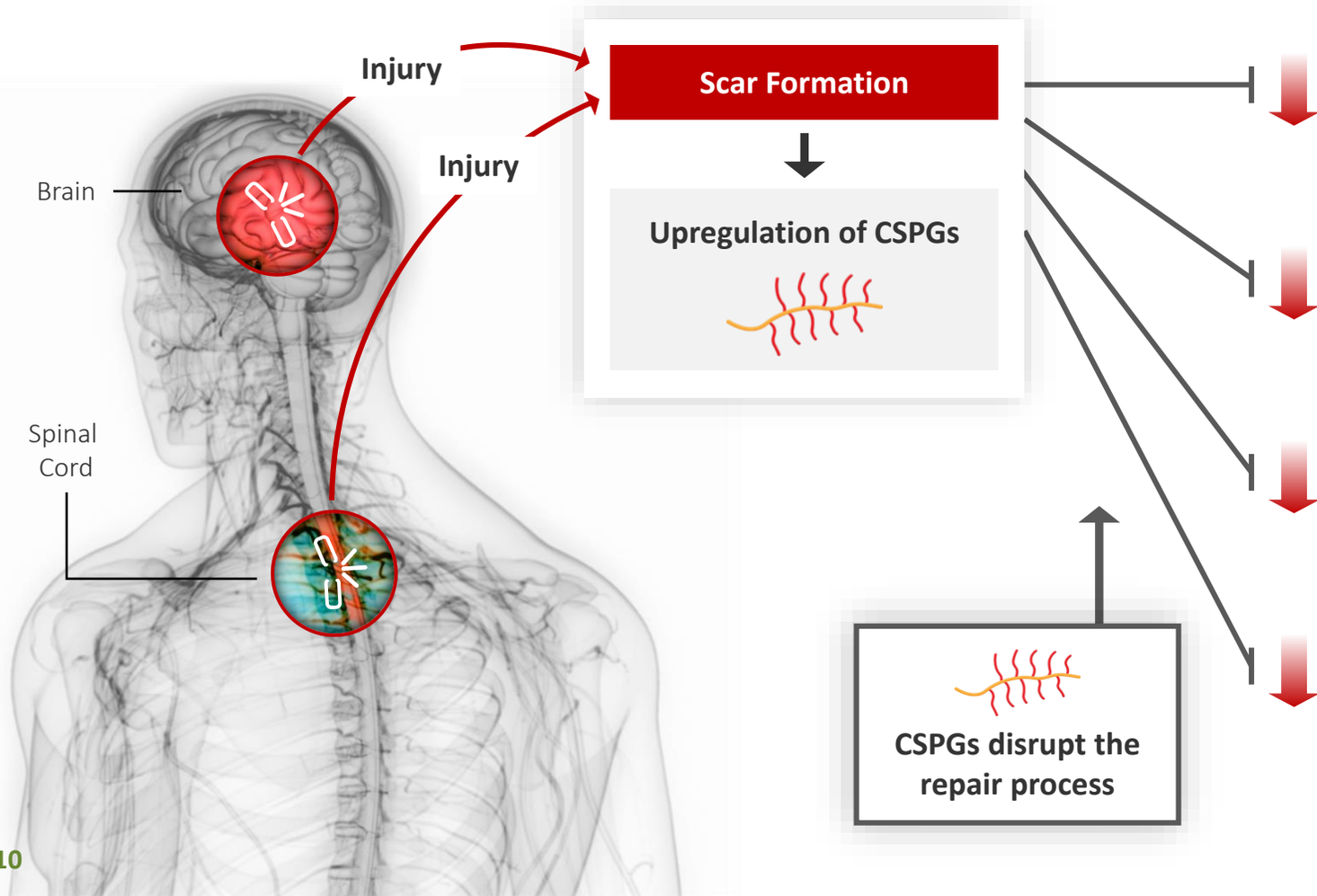
NVG-291 frees neuron from CSPGs and allows it to grow

NVG-291, a 35 amino acid peptide, produced dramatic recovery in a spinal cord injury animal study: the results published in Nature¹ are now cited in over 327 publications

Administered systemically by a **daily subcutaneous injection**

Includes a transporter that **facilitates crossing the blood brain barrier**

The Body's Powerful Innate Repair Mechanisms *Disrupted by CSPGs*



Repair Mechanisms:



Plasticity

The creation of new neuronal connections and rewiring of existing ones



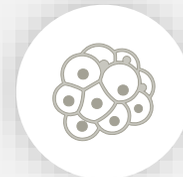
Axonal Regeneration

The ability of a severed axon to reestablish connectivity with other neurons



Remyelination

The process of repairing damaged myelin – the fatty substance that protects axons and enables fast electrochemical transmission

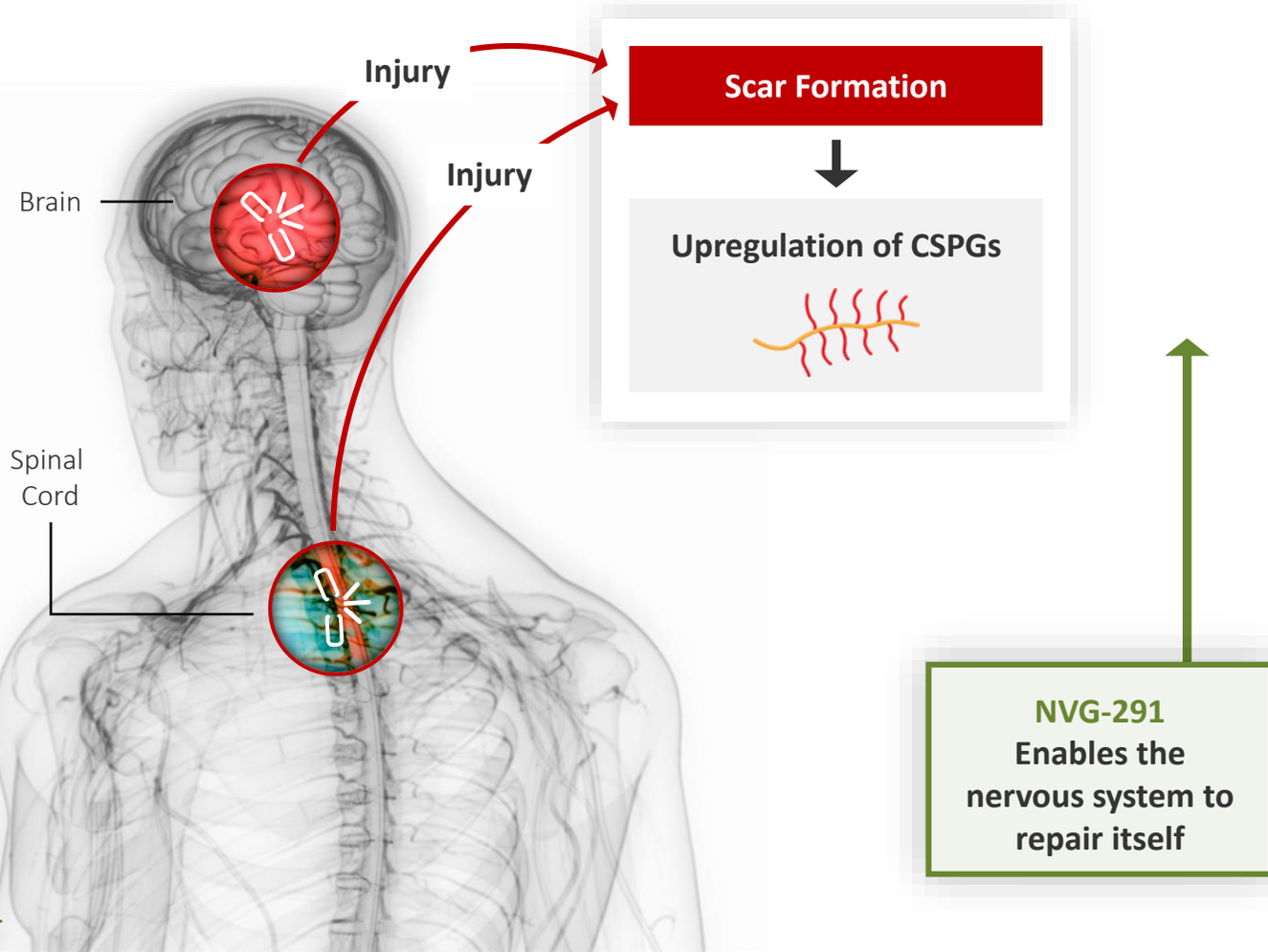


Others

Stem cell preservation/migration
Autophagy
Microglial shifting

NVG-291

Takes the Brakes off Natural Repair Mechanisms



Repair Mechanisms:



Plasticity

The creation of new neuronal connections and rewiring of existing ones



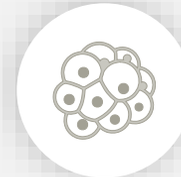
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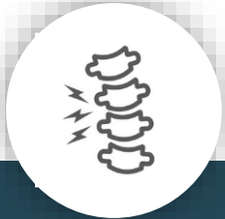
Stem cell preservation/migration
Autophagy
Microglial shifting

NVG-291 Broadly Restores Function

NVG-291 Has Demonstrated Dramatic Repair

in multiple animal models of neurological injury/disease, as documented in 15+ peer-reviewed papers

ACUTE SPINAL CORD INJURY



- Motor
- Sensory
- Bladder

PERIPHERAL NERVE INJURY



- Motor
- Sensory

MULTIPLE SCLEROSIS



- Motor

OPTIC NEURITIS



- Visual

STROKE



- Motor
- Sensory
- Cognition (object recognition)

1. Lang, B.T. et al., *Nature*, 518, 404–408. (2015).
2. Rink, S. et al., *Experimental Neurology*, 309, 148–159. (2018).
3. Ham, T.R. et al., *Ann Biomed Eng*, 47, 744–753. (2019).
4. Ham, T.R. et al., *Materials Science and Engineering: C*, 110, 110656. (2020).

1. Li, H. et al., *Scientific Reports*, 5, 1–14. (2015).
2. Yao, M. et al., *Neuropharmacology*, 144, 208–218. (2019).

1. Luo, F. et al., *Nature Communications*, 9, 1–16. (2018).

1. Niknam, P. et al., *Molecular and Cellular Neuroscience*, 99, 103391. (2019).

1. Unpublished data provided by Dr. Agnes Lou, University of Cincinnati

NVG-291: Potential to Treat All Types of Nervous System Damage

TRAUMA

Acute Spinal Cord Injury

Chronic Spinal Cord Injury

Traumatic Brain Injury

DISEASE

Multiple Sclerosis

Alzheimer's Disease

Stroke

ALS

Frontotemporal Dementia

Parkinson's Disease

NVG-291



NervGen
Priorities

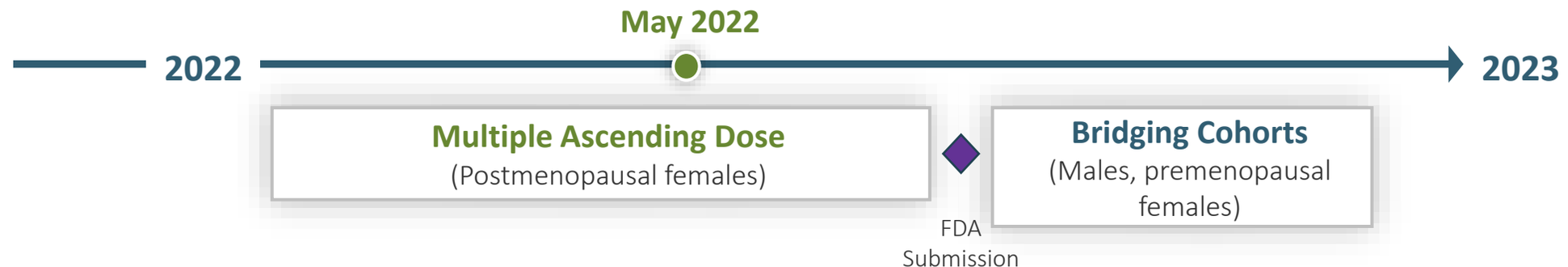
NVG-291 - Phase 1 Clinical Trial in Progress

Single Ascending Dose (SAD) – *COMPLETED*

- NVG-291 was well tolerated at a high dose
 - 170% higher than the equivalent highest dose in preclinical efficacy studies
 - >100x higher than the lowest efficacious dose in studies
- NVG-291 was rapidly distributed in the blood
- The calculated half-life was longer in humans than animals

Multiple Ascending Dose (MAD) – *IN PROGRESS*

- Subjects are dosed once a day for 14 days
- Currently conducting the final cohort
- Dose in second dose cohort was well tolerated and 80% higher than the equivalent highest dose seen in preclinical efficacy studies



Our Phase 1 trial establishes the dose and safety profile necessary for starting all three of our Phase 1b/2 trials



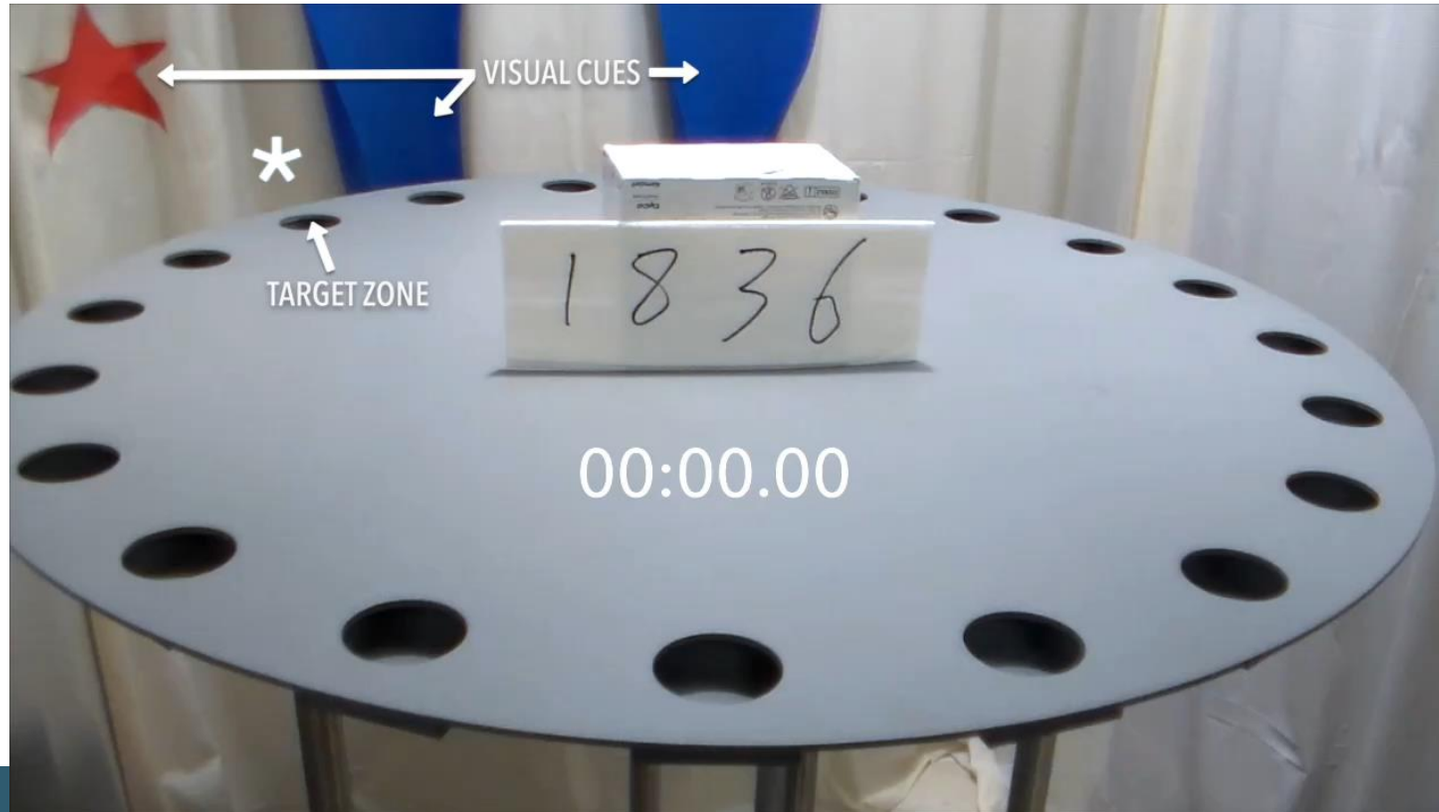
Alzheimer's Disease

No FDA Approved Drug that Results in Sustained Improvement in Cognitive Function

- Symptomatic treatments to improve cognition in Alzheimer's disease are unsatisfactory
- A disease-modifying therapy approved in 2021 has questionable benefit in slowing cognitive decline
- NervGen's goal is to repair damage and improve cognitive function

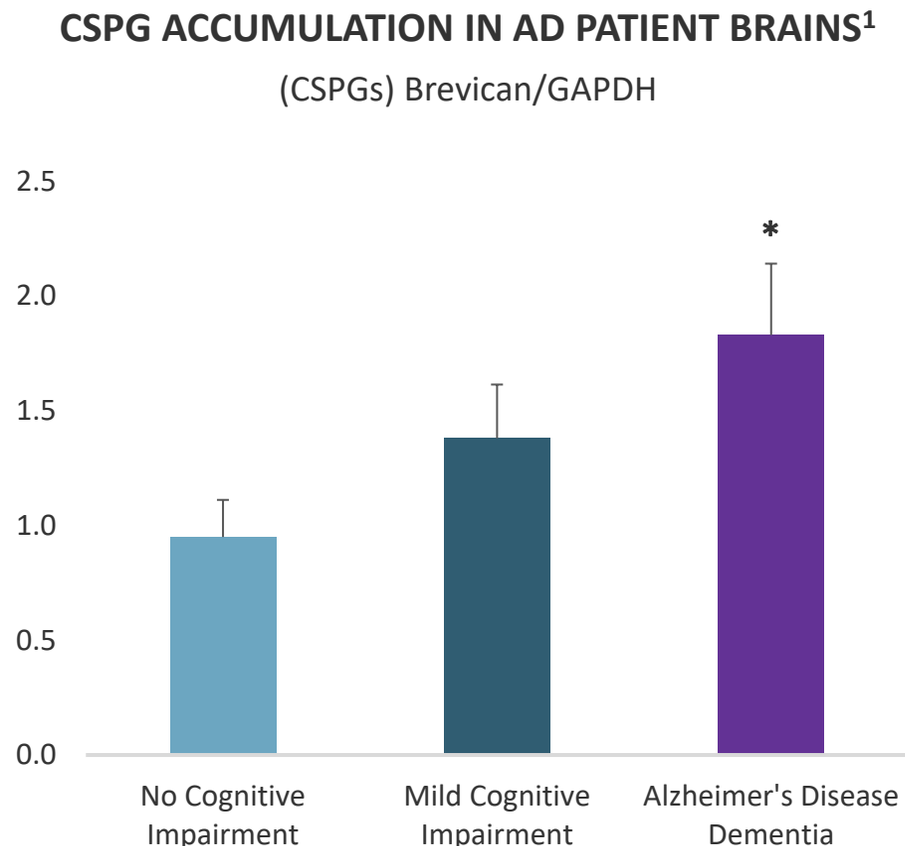
NVG-291 – Improves Memory and Spatial Learning

STROKE MODEL



Significant improvement in cognitive function
when treated 24 hours - and even *7 days* - after a stroke

NVG-291 Pathway to Treat Alzheimer's Disease



* p<0.05 compared to NCI

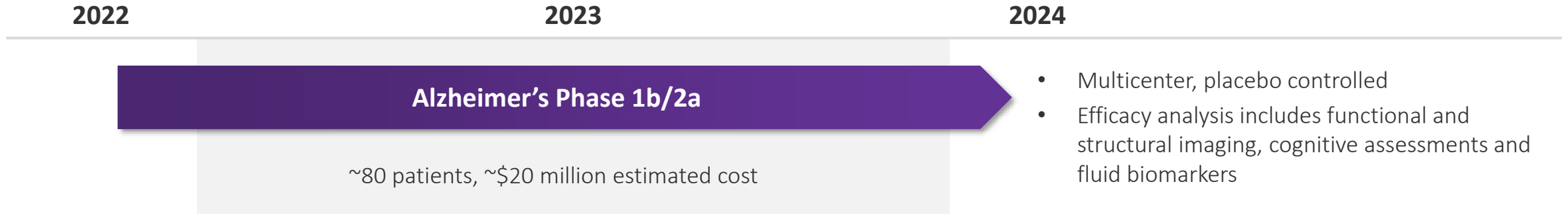
¹ Howell, M.D. et al., Acta Neuropathol Commun, 3, 54. (2015). ² Yang et al., Experimental Neurology (2015). ³ Vegh et al., Acta Neuropathologica Communications (2014). ⁴ Gu et al., BioRxiv (2016)

Preclinical studies have demonstrated that breaking down CSPGs **improves Alzheimer's symptoms^{2,3}**

Removing PTP σ **improves cognitive function** in Alzheimer's models⁴

NVG-291's multiple modes of action, **plasticity, axonal regeneration and remyelination** have the potential to benefit patients suffering from Alzheimer's

NVG-291 Safety/Efficacy Studies in Alzheimer's Disease Patients



World-class Advisory Board with experts in research, clinical design, cognitive assessments and biomarkers

<p>Jeffrey Cummings, MD, ScD University of Nevada</p>	<p>Originator, Neuropsychiatric Inventory (NPI)</p>	<p>Reisa Sperling, MD Harvard Medical School; Massachusetts General Hospital</p>	<p>Led NIA-Alzheimer's Assoc. guideline development group; Serves on National Institute on Aging Advisory Council</p>
<p>Martin Farlow, MD Indiana University School of Medicine</p>	<p>Led/contributed to >230 clinical trials; authored 493 peer reviewed research papers and 509 abstracts</p>	<p>Michael Weiner, MD University of California, San Francisco</p>	<p>Leader in development of MRI and PET for investigating and diagnosing neurodegenerative diseases</p>
<p>Bruce Lamb, PhD Indiana University School of Medicine</p>	<p>World-expert on biological underpinnings of Alzheimer's disease and related dementia</p>	<p>Henrik Zetterberg, MD, PhD University of Gothenburg, University College London</p>	<p>World expert in blood-based biomarkers in neurological disorder</p>
<p>George Perry, PhD University of Texas, San Antonio</p>	<p>Current and founding Editor-in-Chief of the Journal of Alzheimer's Disease</p>		

A man with a beard, wearing a white t-shirt and grey pants, is sitting in a wheelchair. He is looking out a large window on the left side of the frame. The room has a light-colored floor and a white wall. A diagonal beam of light or a structural element is visible in the background.

Spinal Cord Injury

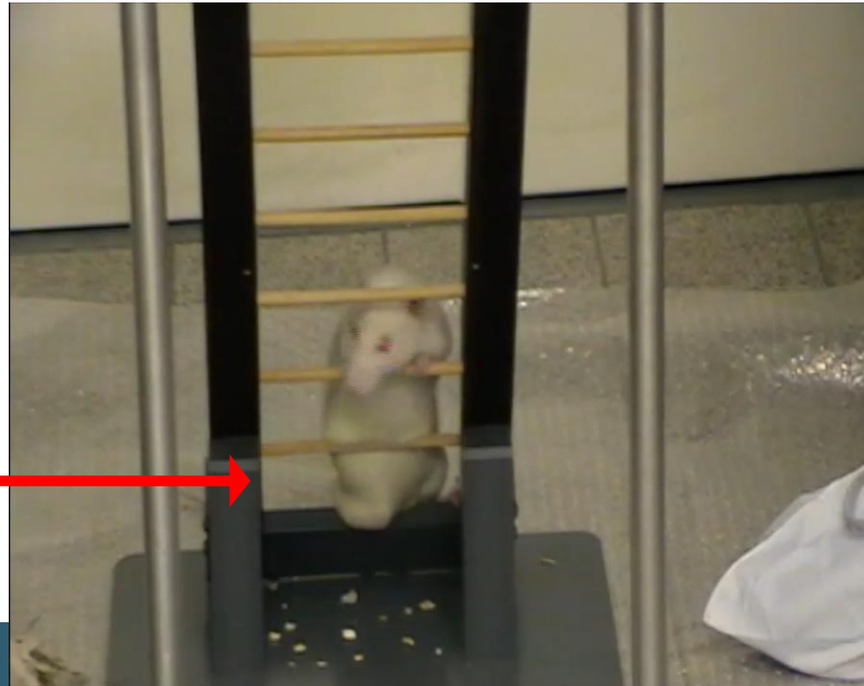
No FDA Approved Drug that Improves Function

- NervGen's goal is to improve motor, bladder/bowel/sexual and/or sensory function

NVG-291 – Dramatically Repairs Spinal Cord Injury

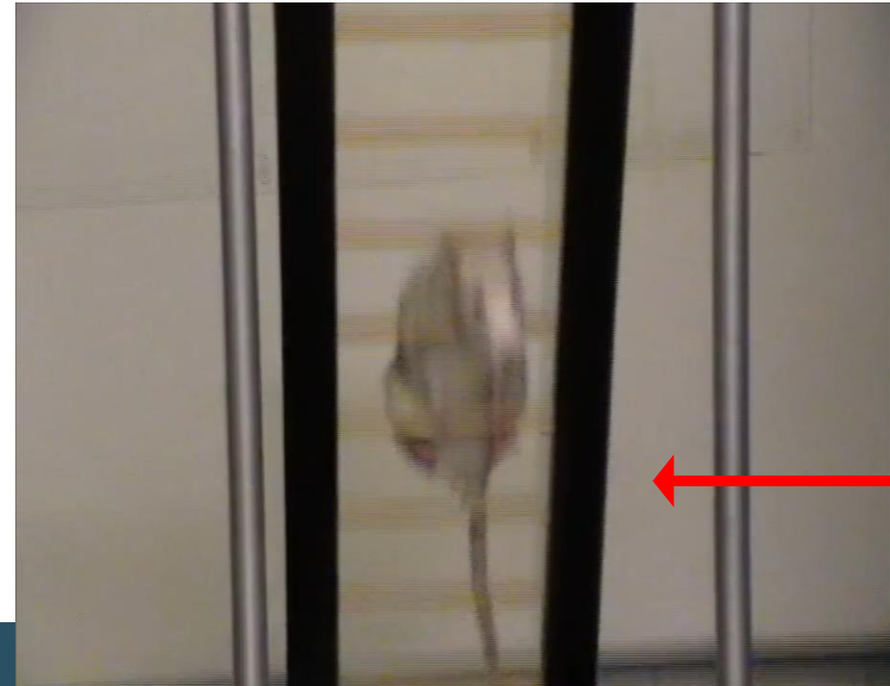
SEVERE SPINAL CORD INJURY MODEL

Representative of Placebo Group



Hind legs are immobile

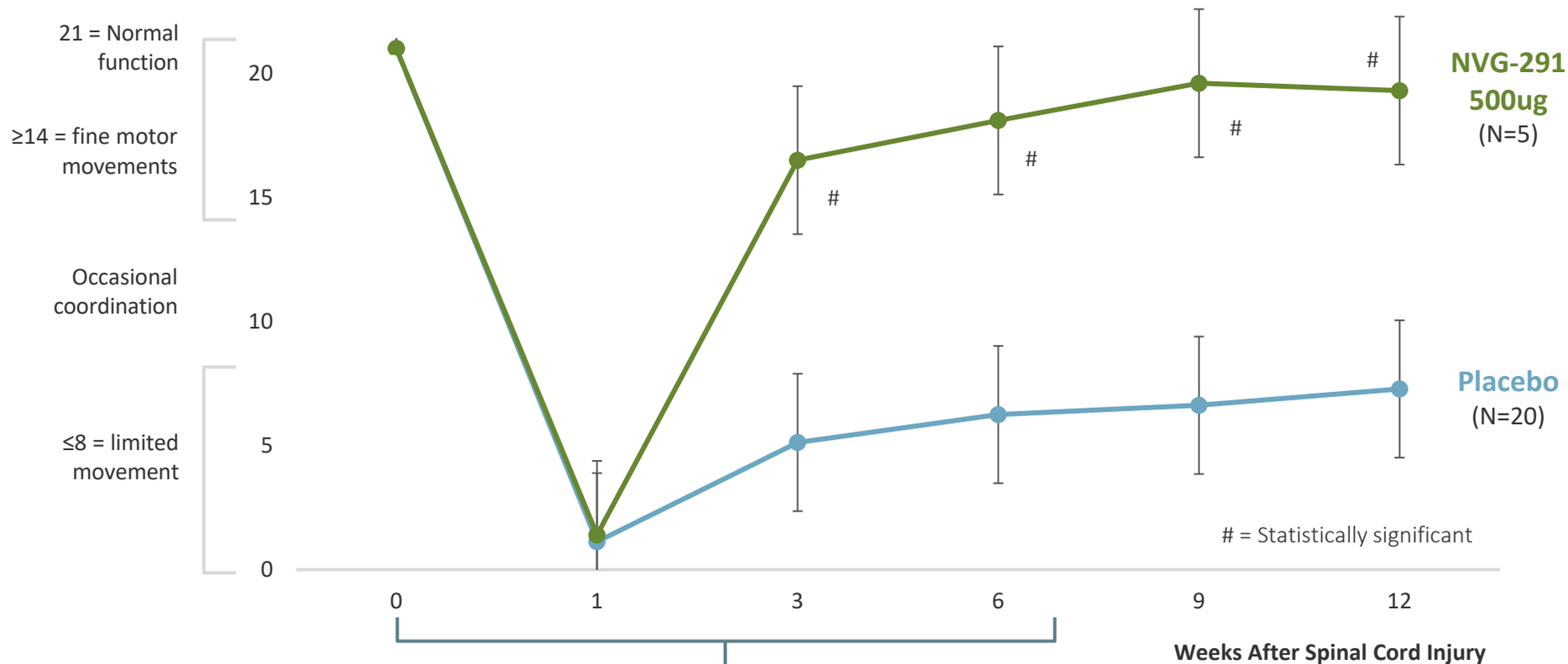
Representative of NVG-291 Group



Significant motor recovery: consistent coordination, toe clearance, tail held high consistently

Spinal Cord Injury – NVG-291 Promotes Functional Recovery

BBB Scale = Standard measure of mobility



Animals treated with once daily systemic subcutaneous injection from 24hrs to 7 weeks post injury

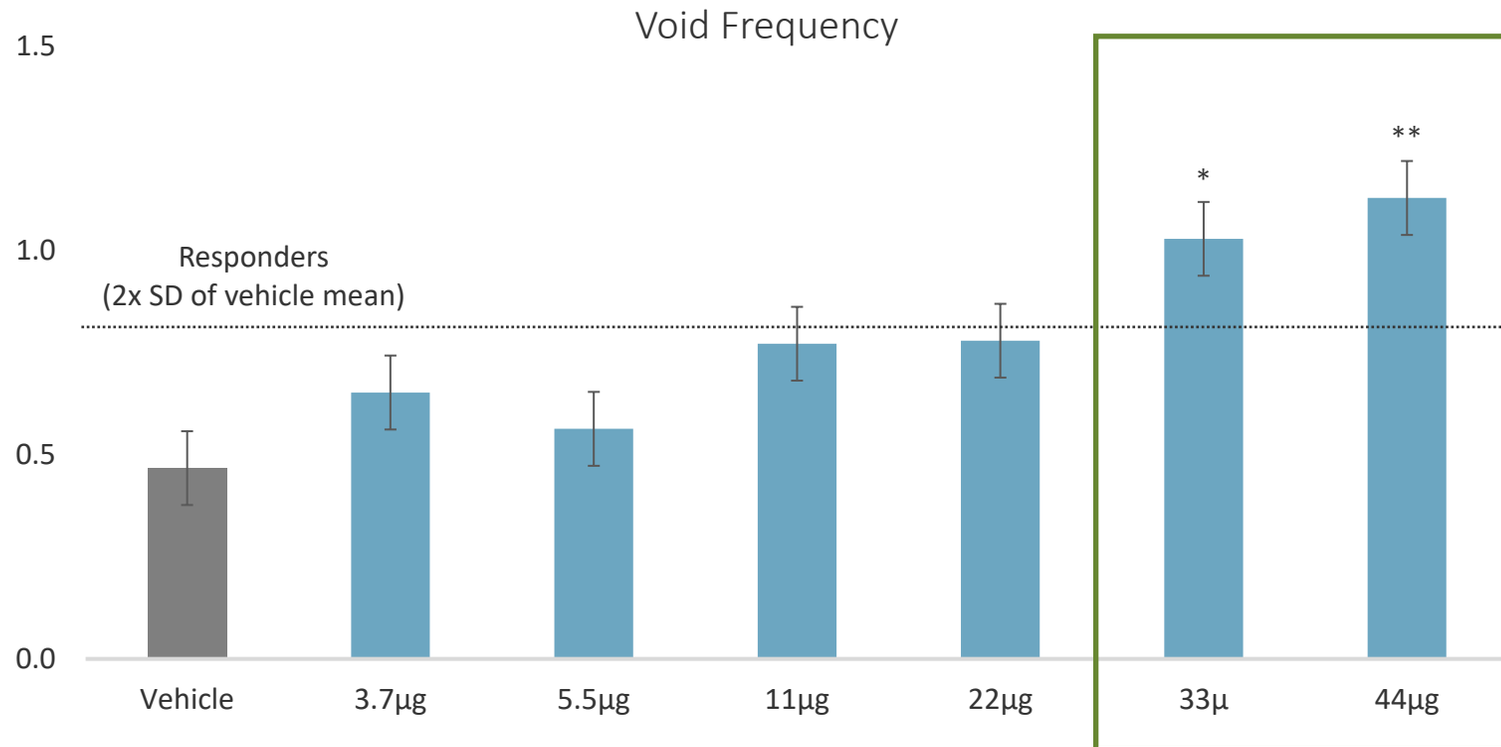
UNPRECEDENTED RESULTS

Extremely high response rate
50%

Almost **complete recovery** in responding animals

NVG-291 Improves Bladder Function

BLADDER DOSE RESPONSE



100% of animals in the two highest dose groups had improved bladder control function

NVG-291 treatment resulted in a dose dependent improvement in bladder function

Bladder function is a key quality of life measure in the paralyzed population

*p <0.05, ** p <0.01

Lang, B.T. et al., Nature 518, 404-408 (2015)



Multiple Sclerosis Repair No FDA Approved Drug that Improves Function

- MS is an autoimmune disease where the immune system attacks myelin in the central nervous system, and over time this results in increasing disability
- Approved disease-modifying drugs modulate the immune system, which can reduce relapses and slow disability progression, but **none** repair damage
- **NervGen's goal is to repair/remyelinate the damage from MS, thereby improving function**

NVG-291 Restores Motor Function in Multiple Sclerosis

Representative of Placebo Group

Score never improves from 3.5



Representative of NVG-291 Group

Score improves to 0.5 in 20 days



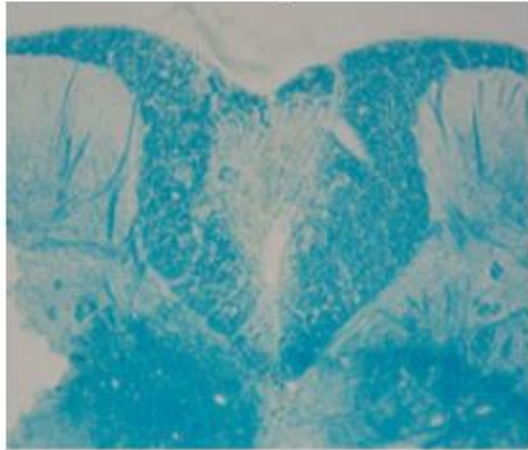
NVG-291 restored motor function in MS model¹,
even when administered after symptoms were fully developed

NVG-291 Remyelinates in Multiple Sclerosis

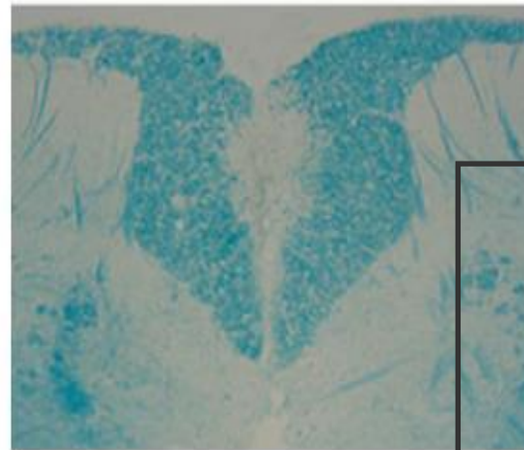
POSITIVE PRECLINICAL RESULTS¹

3 days
post-lesion

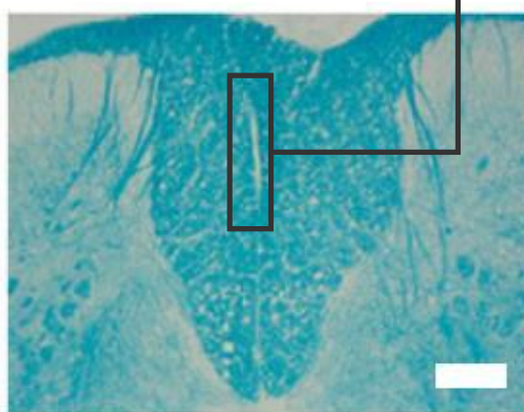
PLACEBO



NVG-291



21 days
post-lesion



Animals treated with NVG-291 had substantially greater remyelination compared to placebo treated animals after 21 days

Histological evidence showed that the desired types of cells were migrating to the lesion, and that these cells were forming myelin

Increased remyelination was accompanied by improvements in nerve conduction

Lesion size in LPC demyelination model

Share and Capital Structure

Exchange/Market: Ticker	TSX: NGEN.V	OTCQX: NGENF
Recent Share Price (May 11, 2022)	CA \$2.17	US \$1.67
Shares Outstanding	47.4 million	
Fully Diluted	63.6 million (~7.1 million options, ~9.1 million warrants)	
Insider Ownership	10%	
~Cash & Cash Equivalents (March 31, 2022)	CA \$12.8 million	US \$10.3 million

Upcoming Value Drivers

ADVANCED CLINICAL TRIAL PROGRAM

	PHASE		INITIATION		READOUT
Alzheimer's Disease	1b/2a		Q4 2022		2024
Spinal Cord Injury	1b/2a		Q4 2022		2023
Multiple Sclerosis	2		Q1 2023		2024

- Phase 1 study topline MAD data (2022)
- Preclinical study results in stroke, chronic spinal cord injury and Alzheimer's disease models which could transform treatment paradigms
- Awarding of privately funded and US Department of Defense sponsored grants (2022)
- Uplisting to Nasdaq (2022)

Investment Highlights

NVG-291 has the potential to **redefine treatment paradigms** for neurological disorders

Improvement demonstrated across 6 different animal models in fine and gross motor control, sensory function, autonomic functions, visual acuity, memory & learning, in many cases **unprecedented**

Pipeline addresses **significant unmet medical needs** in spinal cord injury, multiple sclerosis and Alzheimer's disease

Pipeline addresses **very attractive commercial opportunities**

Experienced management team, board & scientific advisors



Enabling the Nervous System to Repair Itself

 www.nervgen.com

 [@NervgenP](https://twitter.com/NervgenP)

 [NervGen Pharma Corp.](https://www.linkedin.com/company/nervgen-pharma-corp)