



Platform Therapy for Alzheimer's and
Related Neurodegenerative Conditions

Turning Irreversible Into Recoverable

Restoring neurovascular integrity with a first-in-class biologic therapy

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William D. Schwieterman, M.D.
CEO, Stream Biomedical Inc.
251-377-4828 | wds@streambiomed.com

streambiomedical.com

Non-confidential disclosure

For millions living with neurodegenerative disease,
recovery may finally be within reach.



A GLOBAL HEALTH CRISIS

Neurodegenerative diseases impact tens of millions worldwide and remain one of the greatest unmet needs in medicine.



THE LIMITS OF CURRENT APPROACHES

Current approaches primarily target downstream pathology while leaving the underlying neurovascular dysfunction unresolved.



A DIFFERENT KIND OF THERAPY

Stream Biomedical is developing rLG3, a novel recombinant protein designed to restore brain function through **neurovascular repair**.

Failure of neurovascular repair drives **neurodegeneration**

Growing evidence shows that dysfunction of the neurovascular unit is a central driver across neurological diseases.



Pericyte loss and vascular instability



Impaired cerebral perfusion



Blood-brain barrier dysfunction



Chronic neuroinflammation
and impaired repair

Over time, breakdown of these systems
leaves the brain **unable to maintain or repair itself.**



Neurovascular
Restoration



Vascular Integrity
& Repair



Enhanced Perfusion
& Support



Targeted, Systemic
Neuroregeneration

What if the brain could recover?

Stream is developing a new class of biologics — **vascular matrikines** — designed to restore neurovascular integrity by amplifying endogenous repair signaling.



Restores neurovascular stability and repair
Reinforces the brain's capacity to maintain and heal its own vascular network.



Enhances vascular integrity and neuronal support
Improves perfusion and the microenvironment that neurons depend on.



Targets shared upstream mechanisms
Addresses common drivers of neurodegeneration across multiple diseases.

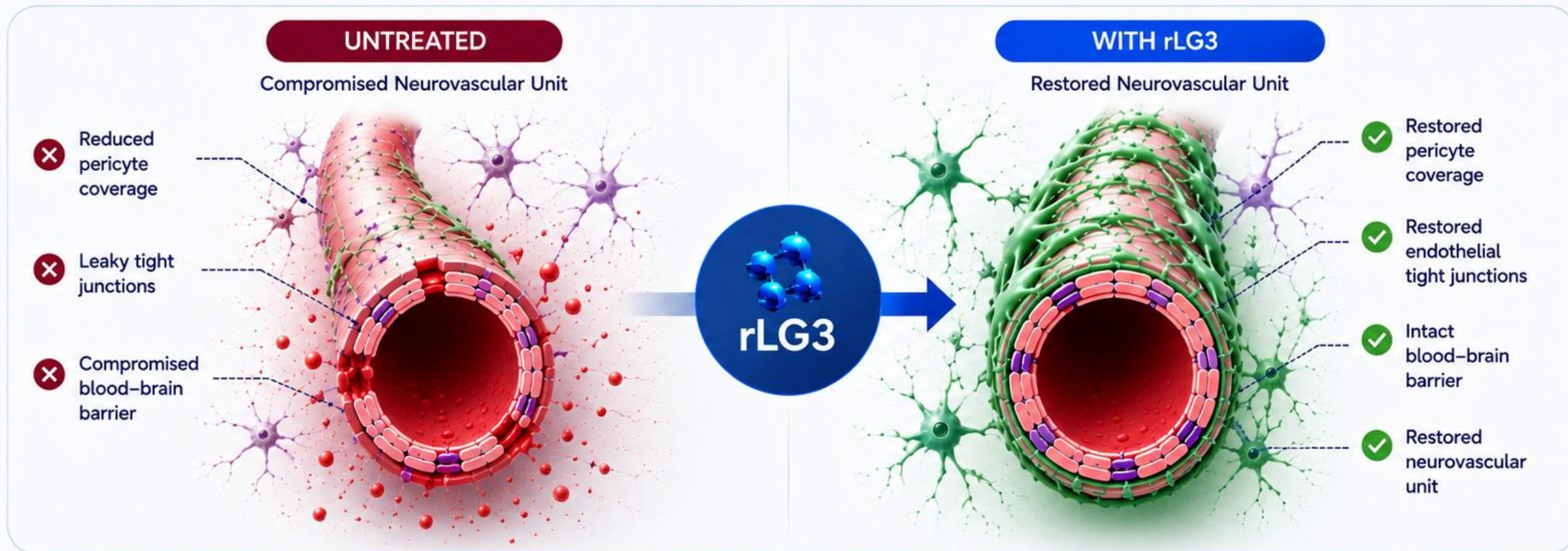


A unified **neurovascular approach to neurodegenerative disease.**

Introducing rLG3: A First-in-Class Recombinant Biologic

Derived from a Natural Endogenous CNS Repair Protein

rLG3 restores neurovascular integrity and CNS homeostasis following injury by restoring pericyte coverage and endothelial tight junctions.



Restoring the brain's own repair mechanisms to **protect, heal and restore.**

rLG3 restores pericyte coverage by engaging integrin receptors.

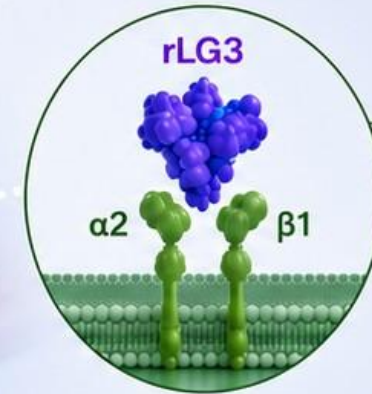
rLG3 is a vascular matrine that binds to integrin receptors on pericytes and endothelial cells to promote pericyte recruitment, adhesion, and stabilization on blood vessels.



The result: Restored pericyte coverage.
Stronger vessels. Healthier brain.

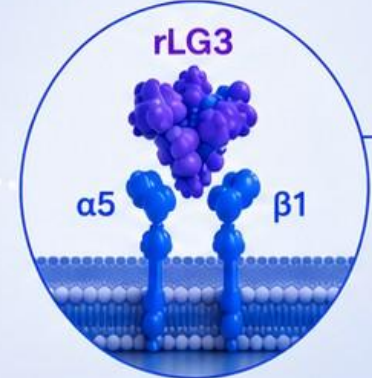
Pericyte

Endothelial cell



rLG3 binds to $\alpha2\beta1$ integrin on pericytes

This interaction promotes pericyte recruitment, adhesion, and stabilization on the vessel wall.



rLG3 binds to $\alpha5\beta1$ integrin on endothelial cells

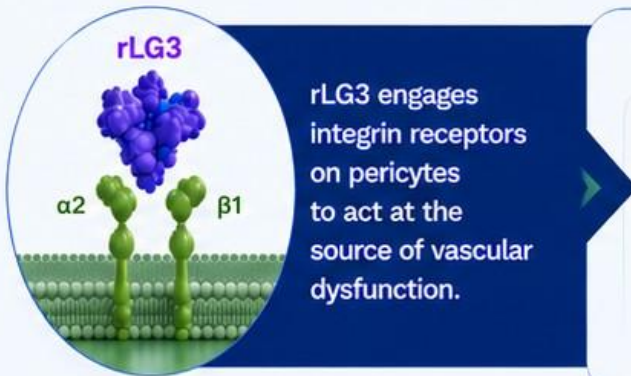
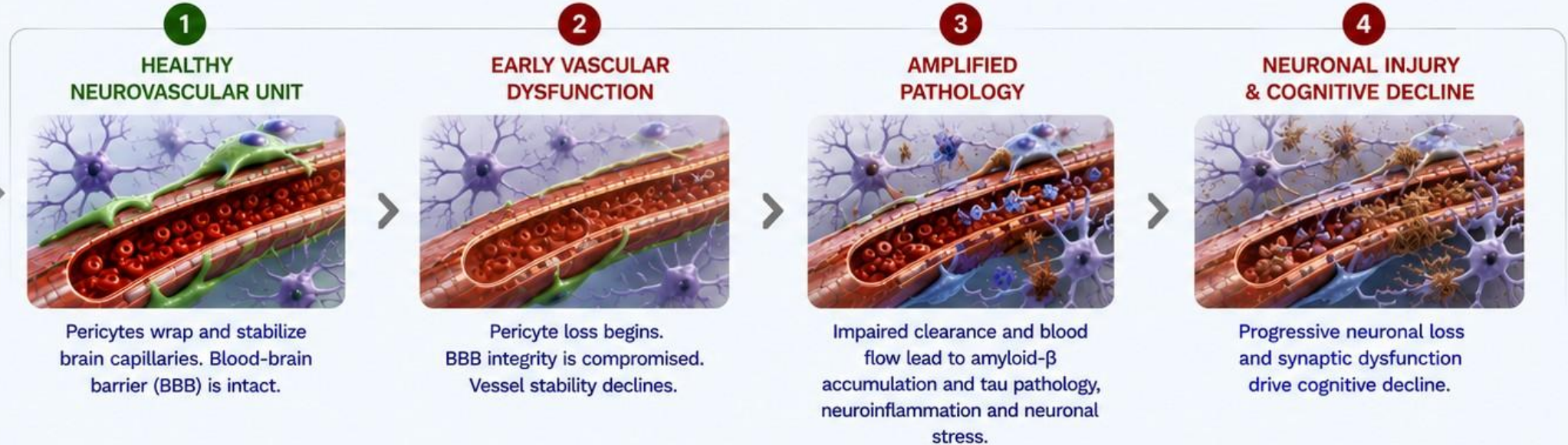
This interaction supports endothelial cell function and vessel integrity.

rLG3 acts **early** to restore the neurovascular foundation

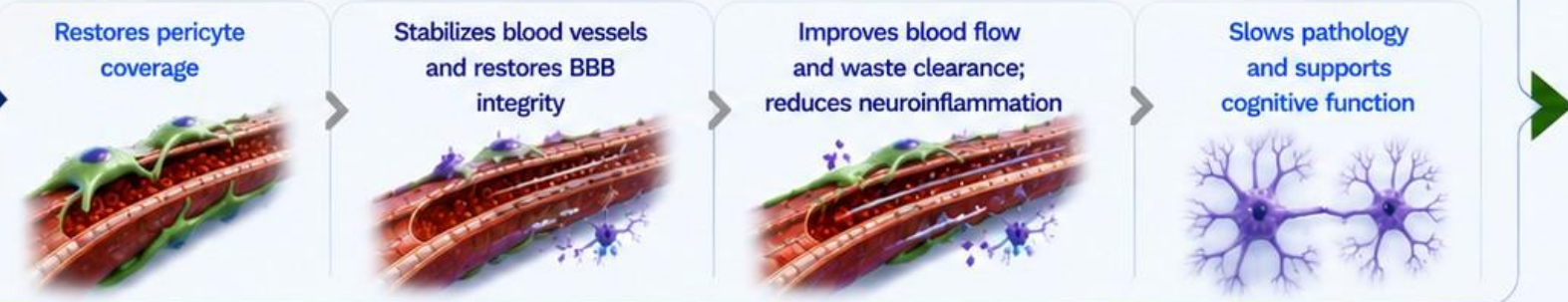
In Alzheimer's disease, vascular dysfunction begins early—driving and accelerating pathology.
rLG3 targets the earliest vascular changes to restore pericyte coverage and support a healthy brain.

THE ALZHEIMER'S DISEASE CASCADE

Vascular dysfunction occurs early—often before clinical symptoms.



— ACTING EARLY. RESTORING VESSELS. PROTECTING THE BRAIN. —



Why Pericytes Matter: Published Human and Preclinical Data Show Neurovascular Dysfunction Drives Downstream Pathology

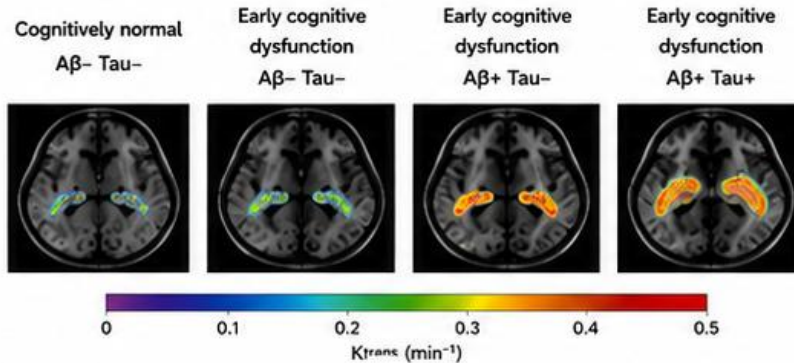
Pericyte injury and BBB breakdown occur early, predict cognitive decline, and cause downstream pathology.

1 HUMAN EVIDENCE

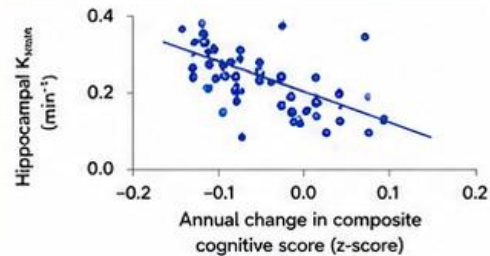
BBB breakdown occurs early and predicts cognitive decline

Nation et al., Nature Medicine, 2019

Hippocampal BBB Permeability (DCE-MRI)



BBB leakage is present in early cognitive dysfunction, before substantial amyloid or tau pathology.



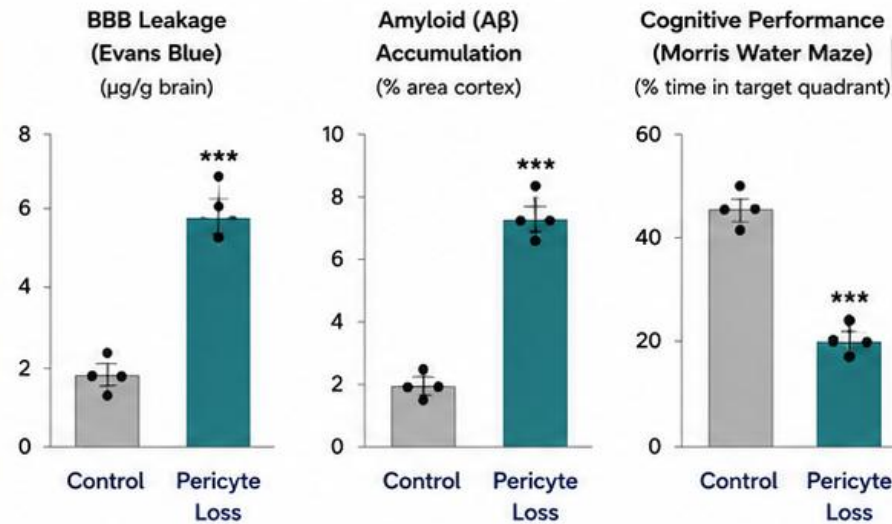
Higher hippocampal BBB permeability independently predicts future cognitive decline ($r = -0.52, p < 0.001$)

2 PRECLINICAL EVIDENCE

Pericyte loss causes BBB breakdown, pathology, and cognitive deficits

Sagare et al., Neuron, 2013

Mouse model: Pericyte depletion in APP mice



Pericyte loss is sufficient to drive BBB breakdown, increased amyloid accumulation, and cognitive impairment.

KEY TAKEAWAYS



Occurs Early

BBB dysfunction and pericyte injury are detectable in early cognitive dysfunction, before amyloid or tau accumulation.



Predicts Decline

Neurovascular dysfunction independently predicts future cognitive decline, even after controlling for amyloid and tau.



Drives Pathology

Pericyte loss causes BBB breakdown, increases amyloid accumulation, and leads to cognitive deficits.

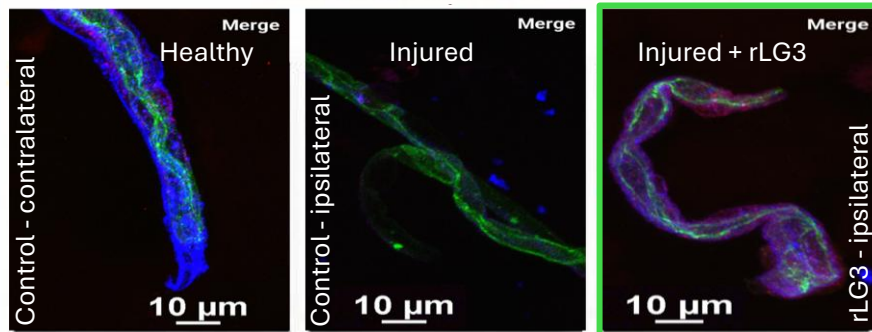


Published human and preclinical studies support pericyte / BBB dysfunction as an upstream driver of neurodegenerative processes.

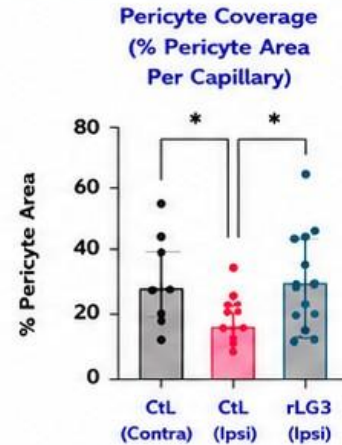
rLG3 Restores Neurovascular Integrity Driving Amyloid Clearance and Cognitive Improvement

1. Restored Pericyte Coverage of Brain Capillaries

Acute Ischemic Stroke Murine Model

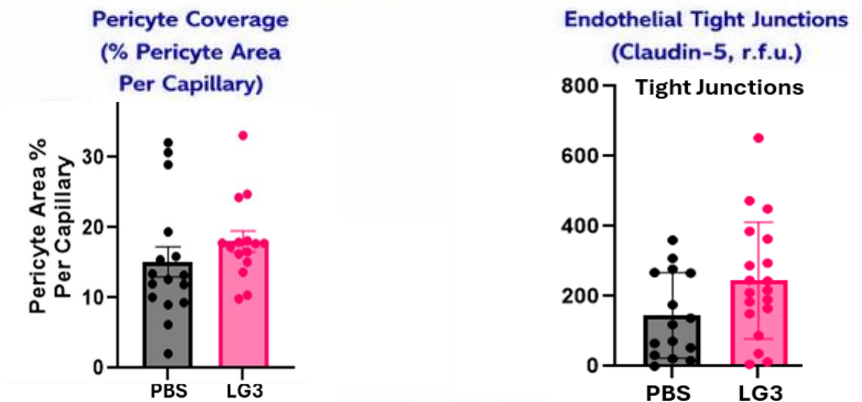


PDGFR-β (Pericyte) Claudin-5 (Endothelial) Lectin (Vessel)



2. Strengthened BBB Integrity

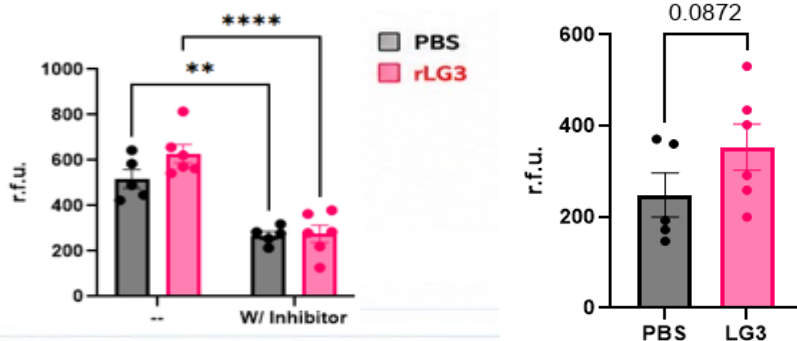
5XFAD Murine Alzheimer's Model



3. Increased Efflux of Amyloid Protein

db/db mice

Amyloid Efflux in Freshly Isolated Brain Capillaries

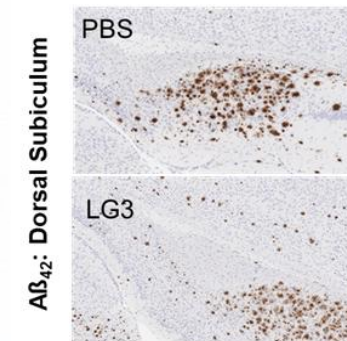


4. Downstream Impact: Reduced Amyloid Pathology

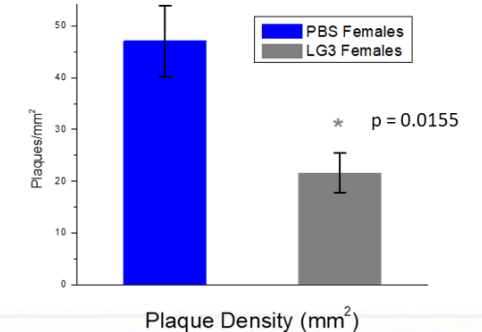
Hippocampus (Main site of AD pathology)



5XFAD Amyloid Model



Amyloid Protein Clearance Amyloid (6E10) Immunofluorescence



rLG3 Treatment
Restores Neurovascular
Health and Function



Restores
pericyte
coverage



Strengthens
BBB
integrity



Increases
amyloid
efflux



Reduces
amyloid
pathology



Supports
cognitive
function

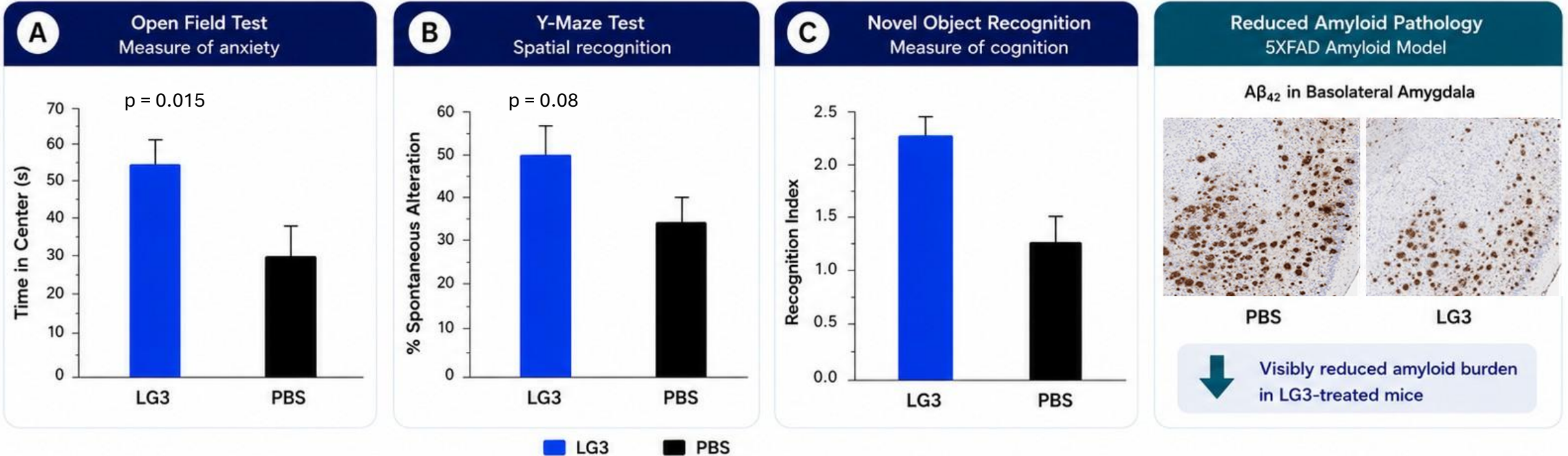


Broad neurovascular
benefits with
downstream impact

rLG3 Improves Cognitive Function in 5xFAD Mice



LG3 treatment significantly improved cognitive and emotional function across multiple behavioral domains, concurrent with **reduced amyloid burden in the basolateral amygdala**.



LG3 increased time in the center of the arena, indicating reduced anxiety



LG3 improved spatial working memory as measured by Y-maze



LG3 enhanced novel object recognition, indicating improved cognition



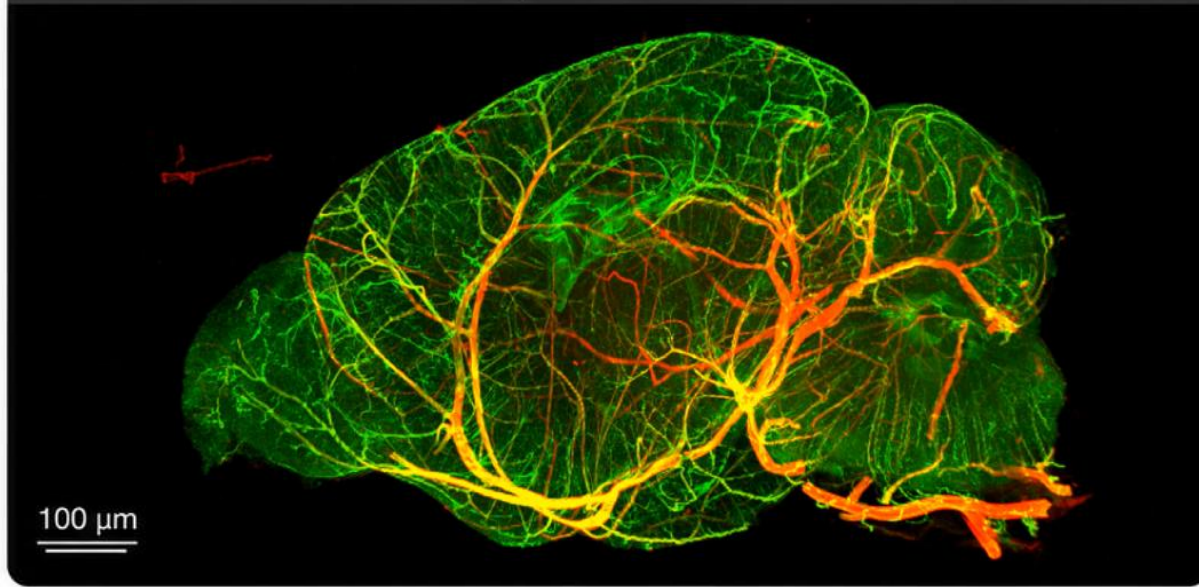
Improvements in behavior coincide with reduced amyloid burden in the basolateral amygdala

n=7-8/group

rLG3 Preserves Cerebrovascular Integrity

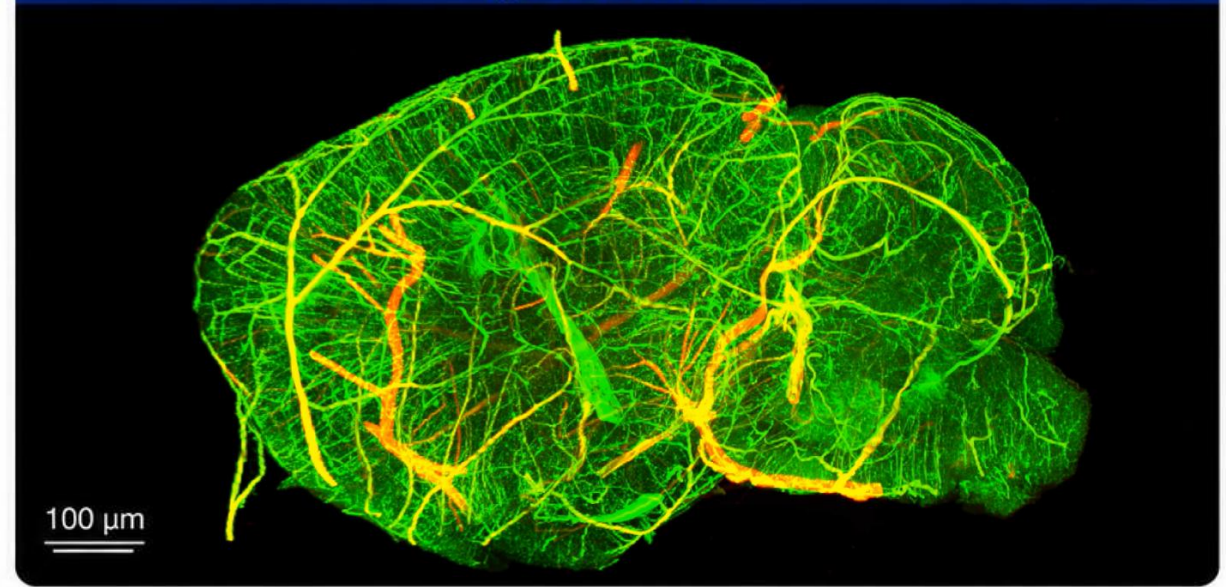
A single dose after reperfusion preserves microvascular structure and density.

Vehicle (Saline)
Injured Brain



Vessel Density: 8.25%
(57% **loss** vs. healthy tissue)

rLG3 Treated
Injured Brain



Vessel Density: 18.92%
(2% **loss** vs. healthy tissue)

Healthy Tissue Vessel Density: 19.19%



rLG3 treatment preserves cerebral microvascular architecture and density, supporting sustained blood flow and neurovascular function.

rLG3: A Multi-Mechanism, Disease-Modifying Therapeutic

Powerful Biology. Broad Potential. Clinic-Ready.



rLG3 PROPERTIES

1



Profound Neuroprotection

2



Eliminates tPA
Hemorrhage Risk

3



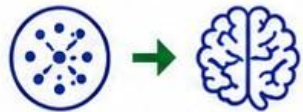
Restores the
Blood Brain Barrier

4



Reduces Biomarkers
of CNS Injury

5



Quickly crosses the BBB
through Active Transport

CLINIC-READY



Recombinant human protein
Manufactured under cGMP



High purity and batch consistency
Robust analytical characterization and QC



Favorable preclinical toxicology
GLP studies complete



Strong intellectual property position
Composition and use of matter patents issued



Clear path to first-in-human trials
IND-enabling package in preparation

rLG3 Phase 1 Trial in Alzheimer's Disease Patients



Biomarker-Focused, Efficient Design to Establish Safety and Demonstrate Target Engagement



OUR FOCUS

Restore vascular integrity in Alzheimer's patients with baseline vascular abnormalities by MRI and fluid biomarkers.



OUR APPROACH

Advanced MRI technology measures vascular function at baseline and 48 hours after LG3 therapy to detect early restoration of vascular integrity.



WHERE WE CONDUCT THIS TRIAL

Leveraging advanced MRI at one of the premier clinical sites in the world.

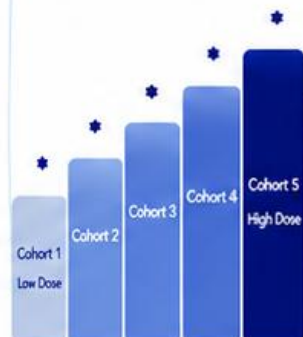


TRIAL SCHEMATIC

Randomized Placebo-Controlled Trial
(3:1 allocation LG3:placebo)

PART 1: SAD Single Ascending Dose

5 Dose Cohorts

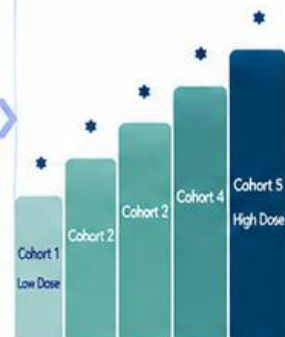


DSMB
Review

- ✓ Safety
- ✓ Tolerability
- ✓ PK/PD
- ✓ Biomarkers

PART 2: MAD Multiple Ascending Dose

5 Dose Cohorts



DOSING & IMAGING PLAN

SHORT, WELL-DEFINED EXPOSURE
3 intravenous doses over 6 days

Day 1



Day 3



Day 6



COMPREHENSIVE ASSESSMENTS

ADVANCED MRI
ASSESSMENTS
Measuring vascular
function



BASELINE
Pre-dose



48 HOURS
AFTER FINAL DOSE

BIOMARKER COLLECTION
Serum & CNS biomarkers



SERUM
Collected at
Baseline
and 48-hour
post-dose



CNS
Collected at
Baseline
and 48-hour
post-dose

Paired imaging and biomarker assessments
detect early restoration of vascular integrity

BIOMARKER READOUTS



VASCULAR (PRIMARY)

- DCE-MRI (Ktrans in hippocampus [BBB leakage])
- Cerebral Blood Flow (ASL MRI)
- Vascular reactivity



CNS INJURY (SECONDARY)

- Serum & CSF biomarkers:
- NfL, GFAP, sTREM2, YKL-40, pTau181, etc.



COGNITION (EXPLORATORY)

- ADAS-Cog13
- CDR-SB
- Memory & executive function tests

SAD = Single Ascending Dose

MAD = Multiple Ascending Dose

DCE-MRI = Dynamic Contrast-Enhanced MRI

ASL MRI = Arterial Spin Labeling MRI

MCI = Mild Cognitive Impairment

rLG3: A Fundamental Solution for Alzheimer's and Other Related Dementias

TURNING IRREVERSIBLE INTO RECOVERABLE

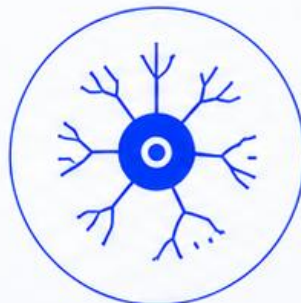


rLG3

A RECOMBINANT
BIOLOGICAL MOLECULE

DERIVED FROM AN
ENDOGENOUS
REPAIR PROTEIN

Harnessing the body's
natural repair pathway
to restore the brain from
the vascular injury.



RESTORES
neurovascular
integrity



REPAIRS
BBB
dysfunction



ENABLES
brain recovery
and resilience

A PROMISING PATH FORWARD

Upstream. Restorative. Clinical-ready.



Appendix

rLG3 Provides Potent Neuroprotection and Restores Function in Traumatic Brain Injury

Single-dose rLG3 reduces cell death, preserves brain tissue, and improves functional recovery.

KEY TAKEAWAYS



POTENT NEUROPROTECTION

Significant reduction in neuronal cell death in the cortex after TBI.



FUNCTIONAL RECOVERY

Rapid and sustained improvements in motor function and grip strength.



BBB INTEGRITY

Marked reduction of Evan's Blue leakage in the impact region.



SINGLE DOSE EFFICACY

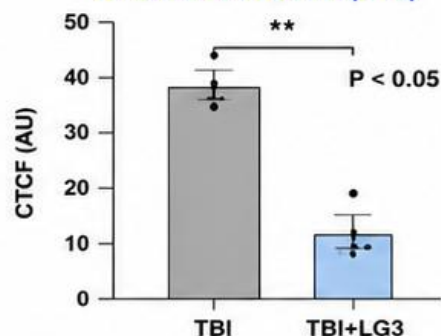
Robust benefits achieved with a single administration of rLG3.



These data demonstrate that rLG3 protects brain tissue and restores function after TBI by preserving vascular integrity and reducing secondary injury.

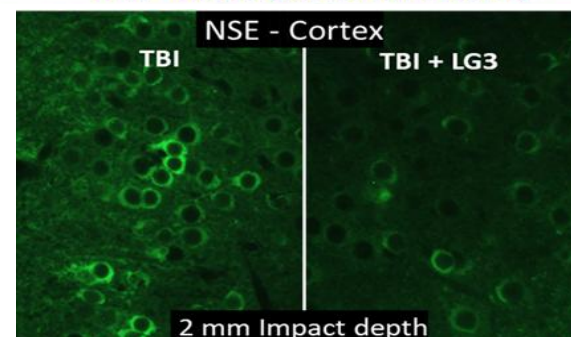
1 REDUCES CELL DEATH

Neuronal Cell Death (NSE)



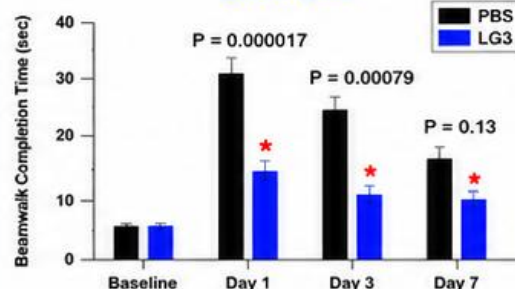
2 PRESERVES CORTICAL TISSUE

NSE – Cortex (Green Fluorescence)

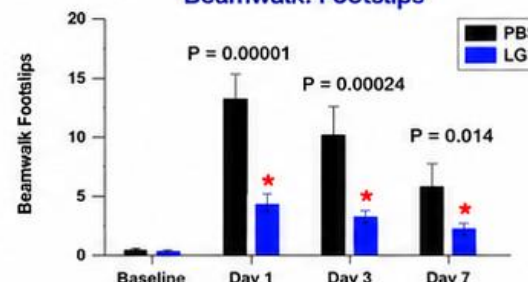


3 IMPROVES FUNCTIONAL PERFORMANCE

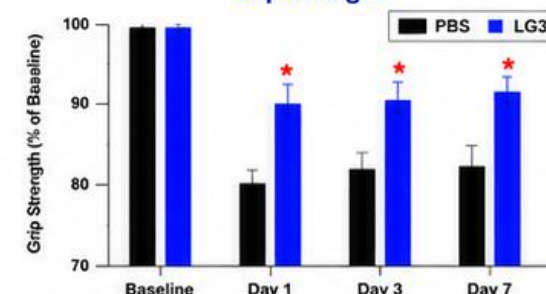
Beam Walk



Beamwalk: Footslips



Grip Strength



4 IMPROVES BBB INTEGRITY

Evan's Blue BBB Leakage (Impact Region)

TBI – Control (PBS)



TBI – LG3

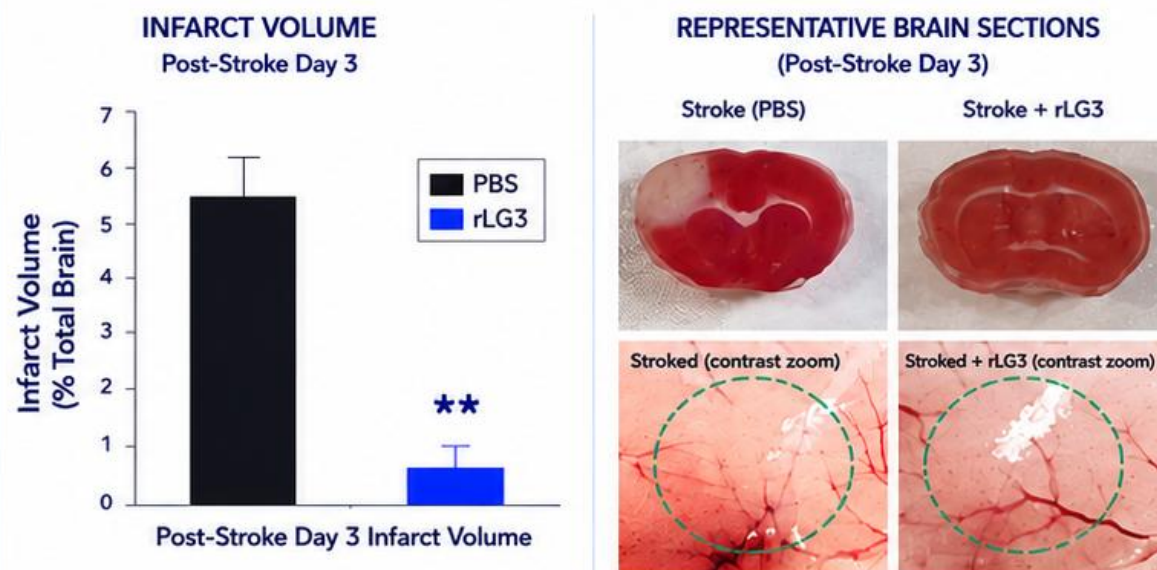


rLG3 Delivers Potent Neuroprotection and Eliminates tPA-Associated Hemorrhage

Restoring vascular integrity improves outcomes and broadens therapeutic potential.

1 ISCHEMIC STROKE MODEL (MCAO)

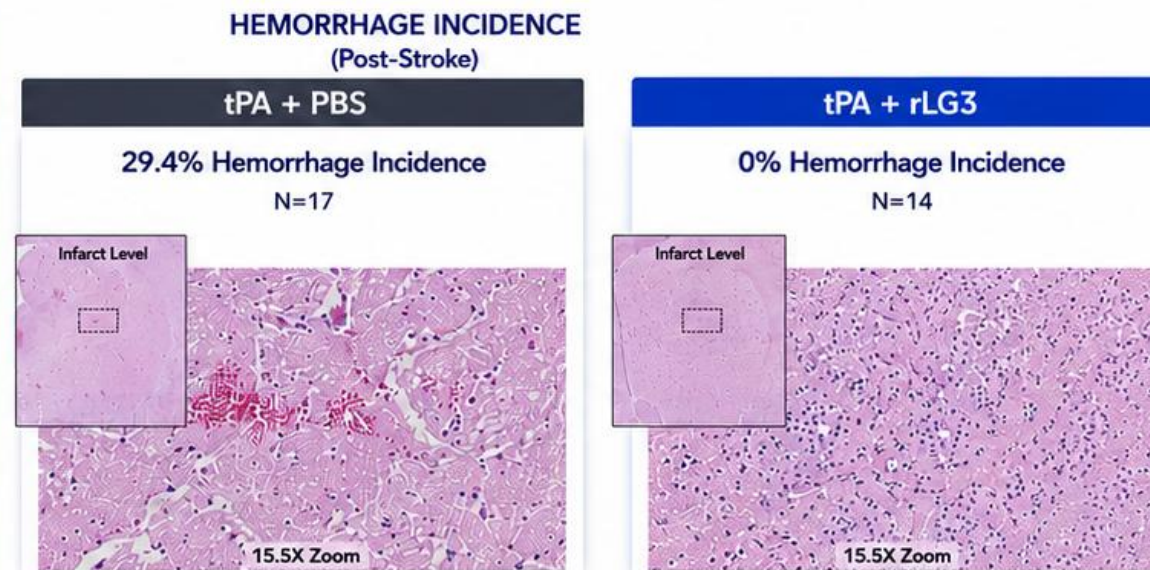
Profound Neuroprotection with Single Dose rLG3



rLG3 markedly reduces infarct volume and preserves brain tissue following ischemic stroke.

2 tPA MODEL OF ACUTE STROKE

rLG3 Eliminates tPA-Associated Hemorrhage



rLG3 completely prevents tPA-associated hemorrhage, supporting vascular protection and enabling safer reperfusion therapy.



Across ischemic stroke models, rLG3 protects brain tissue and preserves vascular integrity—supporting its potential as a disease-modifying therapy for stroke.

The Molecular Genesis of Endogenous LG3

A Conserved, Injury-Induced Response for Neuroprotection and Repair

Endogenous LG3 is generated through a tightly regulated, injury-induced proteolytic process that liberates the LG3 domain from Perlecan Domain V and delivers it to sites of CNS injury where it binds to cell surface integrin receptors to initiate neuroprotective and repair pathways.

1 CNS INJURY



Traumatic or ischemic injury triggers a rapid biological response in the CNS.

2 ACTIVATION OF PLASMINOGEN

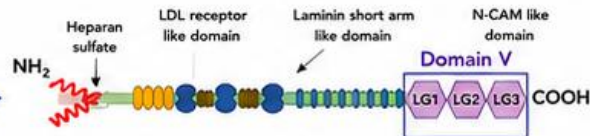
tPA/uPA

- Injury induces release of tPA and/or uPA.
- These serine proteases convert plasminogen to plasmin.



3

CLEAVAGE OF PERLECAN (DOMAIN V)



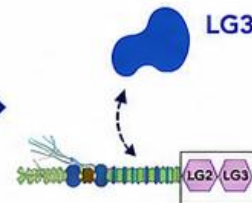
Perlecan (HSPG2)

- Perlecan is a large heparan sulfate proteoglycan in the extracellular matrix and basement membranes.
- Domain V at the C-terminus contains three laminin G (LG) modules (LG1, LG2, LG3).
- Plasmin, caspases and other proteases cleave perlecan within Domain V, releasing the LG3 domain.

Source of LG3:
Perlecan Domain V (C-terminus)

4

RELEASE OF LG3 DOMAIN



- The LG3 domain is liberated from perlecan as a soluble glycoprotein (~19 kDa).
- This is the endogenous form of LG3.

Endogenous LG3
(~19 kDa)

5

HOMING TO THE INJURY SITE



- LG3 homes to the site of injury by binding to upregulated integrin receptors on cells.

Targeted Localization
to Injury Site

6

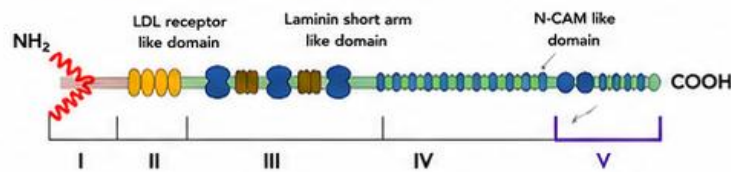
NEUROPROTECTION & REPAIR



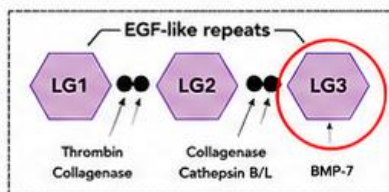
- LG3 binding initiates potent neuroprotective and repair pathways.
- Supports cell survival, BBB integrity, and functional recovery.

Neuroprotection
& Repair

PERLECAN DOMAIN V (SOURCE OF LG3)

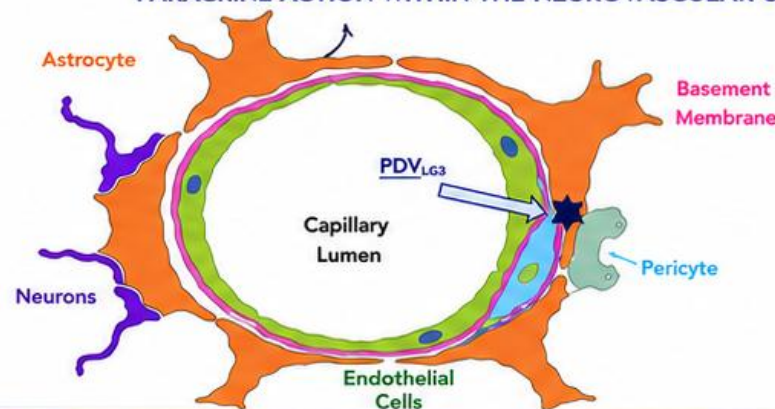


Domain V
Structure
(Endorepellin)



Plasmin, caspases and other proteases cleave perlecan within Domain V, releasing the LG3 domain.

PARACRINE ACTION WITHIN THE NEUROVASCULAR UNIT



Endogenous LG3 acts as a paracrine signal (~40 nm distance) between cells of the neurovascular unit to coordinate survival, inflammation control, and repair.



This is how endogenous LG3 is naturally generated and functions.

rLG3 therapeutically amplifies this intrinsic protective system.