



DENALI

Corporate Overview

October 2023

Disclaimers

Forward-Looking Statements. This presentation contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. These forward-looking statements do not relate strictly to historical or current facts and they may be accompanied by such words as “anticipate,” “believe,” “could,” “estimate,” “expected,” “forecast,” “intend,” “may,” “plan,” “potential,” “possible,” “future,” “will” and other words and terms of similar meaning. All statements other than statements of historical facts contained in this presentation, including, without limitation, statements regarding future results of operations and financial position of Denali Therapeutics Inc. (“Denali” or the “Company”); Denali’s business strategy and business plans, expected progress and expansion, and expected key milestones for Denali’s therapeutic portfolio in 2023 and beyond; Denali’s ability to execute on its tailored commercial strategies and accelerate commercial launch readiness in key markets, including the US and China; expectations relating to the prevalence and potential for Denali’s product candidates to treat various neurodegenerative diseases including MPS I, MPS II (Hunter Syndrome), MPS IIIA (Sanfilippo Syndrome), ALS, MS, PD, AD, FTD-GRN, CLE, UC, and related peripheral inflammatory diseases; expectations and timelines related to planned preclinical studies and clinical trials and the expectations regarding the timing and availability of results and data from such studies and trials; plans, timelines, expectations, and current and future therapeutic and commercial opportunities related to Denali’s Transport Vehicle (TV) platform, including its Enzyme Transport Vehicle (ETV), Antibody Transport Vehicle (ATV), Protein Transport Vehicle (PTV), and Oligonucleotide (OTV) technologies, and other programs enabled by these platforms, as well as potential targets, therapeutic areas, and differentiation strategies; plans, timelines, and expectations relating to DNL310, including safety profile and exploratory clinical outcomes data from the ongoing Phase 1/2 study, enrollment in the Phase 2/3 COMPASS study, and planned regulatory filings and registration potential; plans, timelines and expectations related to DNL126, including its therapeutic potential and the timing of recruitment activities for the planned Ph 1/2 study; Denali’s and Takeda’s plans and expectations regarding DNL593, including enrollment in the ongoing Ph 1/2 Part B trial; expectations and potential benefits relating to ATV:Abeta for the potential treatment of AD; expectations and timelines related to OTV-enabled programs, including their therapeutic and registrational potential; plans, timelines, and expectations relating to the Biogen-led development of BII122/DNL151, including for the Phase 2b trial, as well as other LRRK2 inhibitor molecules; plans, timelines, and expectations related to DNL343, including the timing and availability of data from the ongoing Healey trial and success of the selected model; Denali’s and Sanofi’s plans, timelines, and expectations related to DNL788 and DNL758, including with respect to the availability of data and recruitment of patients for current trials and potential completion dates; the potential benefits and results of the collaborations with Denali’s partners, including Biogen, Sanofi, and Takeda, and the amounts and likelihood of receipt of milestone payments; plans and expectations regarding Denali’s global organization, including the expansion of its medical affairs and clinical operations, the growth of its in-house clinical manufacturing capabilities, and the expected timing and likelihood of success of its commercial growth; and timing and expectations regarding potential additional BBB transporters; are forward-looking statements. Denali has based these forward-looking statements largely on its current expectations and projections about future events.

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OUR PURPOSE: **DEFEAT DEGENERATION**



**RARE NEURODEGENERATIVE
DISEASES**



**AMYOTROPHIC LATERAL
SCLEROSIS (ALS)**



**PARKINSON'S
DISEASE**



**ALZHEIMER'S
DISEASE**



Denali

The name captures the formidable challenges in fighting neurodegenerative diseases but also the unprecedented opportunities enabled by new scientific insights and technologies. With a relentlessly committed team and rigorous effort, breakthroughs appear to be within reach.

OUR PRIORITIES

1 Clinical Execution

- 4 late-stage programs enrolling studies in MPS II, 2x ALS, and PD
- Multiple earlier-stage trials designed for biomarker PoC
- Expansion of clinical operations and medical affairs in Europe
- Building out clinical manufacturing capabilities

2 TV Expansion

- Clinical data from 3 TV-platform enabled programs
- Fourth TV-enabled program advancing towards clinical testing
- Selected OTV targets provides broad range of opportunities
- Expand TV platform potential with additional BBB transporter

3 Commercial Readiness

- Define go-to-market strategies in the US and key global markets
- Outreach to patients and communities in MPS II and ALS to understand unmet needs
- Establish critical medical affairs and commercial capabilities to prepare for early filing scenarios

TV=Transport Vehicle; OTV=Oligonucleotide Transport Vehicle; MPS= mucopolysaccharidoses; ALS=amyotrophic lateral sclerosis; PD=Parkinson's disease; PoC=proof of concept

\$1.19B in cash and investments (as of 6/30/23)

OUR FOCUS AND STRATEGIC PRINCIPLES

OUR FOCUS

Defeat Degeneration



Lysosomal Storage Diseases



Rare Neurodegenerative Diseases (ALS, FTD)



Parkinson's Disease



Alzheimer's Disease

OUR SCIENTIFIC PRINCIPLES

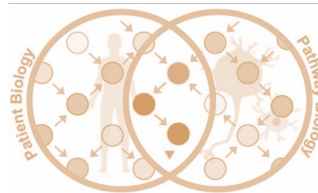
Increase Likelihood of Success



Degenogene Pathways



Brain Delivery



Biomarker-Driven Development

OUR BUSINESS PRINCIPLES

Create Value



Broad Portfolio



Integrated Global Capabilities



Strategic Partnering

OUR SCIENCE: BBB PLATFORMS AND DEGENOGENE PATHWAYS

Published May 27, 2020

SCIENCE TRANSLATIONAL MEDICINE | RESEARCH ARTICLE
BLOOD-BRAIN BARRIER

Brain delivery of therapeutic proteins using an Fc fragment blood-brain barrier transport vehicle in mice and monkeys

Cell

Published Sept 2, 2021

CellPress

Article

Rescue of a lysosomal storage disorder caused by *Gm* loss of function with a brain penetrant progranulin biologic

CellPress

Neuron

Article

Published March 4, 2020

TREM2 Regulates Microglial Cholesterol Metabolism upon Chronic Phagocytic Challenge

Published May 27, 2020

SCIENCE TRANSLATIONAL MEDICINE | RESEARCH ARTICLE
BLOOD-BRAIN BARRIER

Brain delivery and activity of a lysosomal enzyme using a blood-brain barrier transport vehicle in mice

International Journal of
Molecular Sciences

Published July 22, 2020

MDPI

Article

Characterization of Fluid Biomarkers Reveals Lysosome Dysfunction and Neurodegeneration in Neuronopathic MPS II Patients

CellPress

Neuron

Article

Published October 22, 2020

Small-Molecule Modulation of TDP-43 Recruitment to Stress Granules Prevents Persistent TDP-43 Accumulation in ALS/FTD

Published April 8, 2020

NATURE REVIEWS | DRUG DISCOVERY

Leveraging preclinical models for the development of Alzheimer disease therapeutics

International Journal of
Molecular Sciences

Published July 30, 2020

MDPI

Article

High-Throughput Liquid Chromatography–Tandem Mass Spectrometry Quantification of Glycosaminoglycans as Biomarkers of Mucopolysaccharidosis II

nature
neuroscience

Published June 8, 2020

Alzheimer's-associated PLC γ 2 is a signaling node required for both TREM2 function and the inflammatory response in human microglia

Neuron

Published Sept 4, 2019

CellPress

Review

Emerging Microglia Biology Defines Novel Therapeutic Approaches for Alzheimer's Disease

JCI insight

Published October 8, 2021

Iduronate-2-sulfatase transport vehicle rescues behavioral and skeletal phenotypes in a mouse model of Hunter syndrome

Published June 8, 2022





SCIENCE TRANSLATIONAL MEDICINE | RESEARCH ARTICLE

PARKINSON'S DISEASE

Preclinical and clinical evaluation of the LRRK2 inhibitor DNL201 for Parkinson's disease

Denali scientists have generated more than 20 publications and 90 granted patents worldwide

OUR DEVELOPMENT PORTFOLIO

MODALITY	TARGET	BIOLOGY	DRUG CANDIDATE*	DISEASE INDICATION	DEVELOPMENT STAGE				PARTNER
					IND-Enabling	Early	Mid	Late	
LARGE MOLECULE (TV-ENABLED)	Iduronate 2-Sulfatase	Lysosomal Function	DNL310 (ETV:IDS)	MPS II (Hunter)					
	PGRN	Lysosomal Function	TAK-594/DNL593 (PTV:PGRN)	Frontotemporal Dementia-Granulin (FTD-GRN)					
	Sulfamidase	Lysosomal Function	DNL126 (ETV:SGSH)	MPS IIIA (Sanfilippo)					
	Alpha-L-iduronidase	Lysosomal Function	DNL622 (ETV:IDUA)	MPS I (Hurler)					
	Multiple	Multiple	OTV:Multiple	Multiple					
SMALL MOLECULE	LRRK2	Lysosomal Function	BIIB122/DNL151 (LRRK2 inhibitor)	Parkinson’s Disease					
	RIPK1 (CNS)	Glial Biology	SAR443820/DNL788 (RIPK1 inhibitor)	Amyotrophic Lateral Sclerosis (ALS)					
				Multiple Sclerosis (MS)					
	RIPK1 (Peripheral)	Other	SAR443122/DNL758 (RIPK1 inhibitor)	Ulcerative Colitis (UC)					
	eIF2B	Cellular Homeostasis	DNL343 (eIF2B activator)	Amyotrophic Lateral Sclerosis (ALS)					

Biotherapeutics

Small Molecules

 50/50 US Commercial

 Royalty

*Investigational – not approved for treatment

Broad, diverse, and differentiated portfolio, including multiple TV-enabled and small molecule programs in discovery

OUR STRATEGIC PARTNERSHIPS

CO-DEVELOPMENT & CO-COMMERCIALIZATION PARTNERSHIPS



- **LRRK2** inhibitor for Parkinson's and **ATV:Abeta**
- \$1.025B upfront (cash/equity) and \$2B in milestones
- LRRK2: 50/50 profit share in US, 40/60 in China



- **RIPK1** inhibitors for neurological and peripheral inflammatory indications
- \$125M upfront and \$1.1B in milestones
- 50/50 profit share in US/China (CNS)



- **PTV:PGRN** and **ATV:TREM2**
- \$150M upfront (cash/equity) and \$1B in milestones
- 50/50 profit share worldwide

\$1.3B

Total upfront payments¹

>\$3B

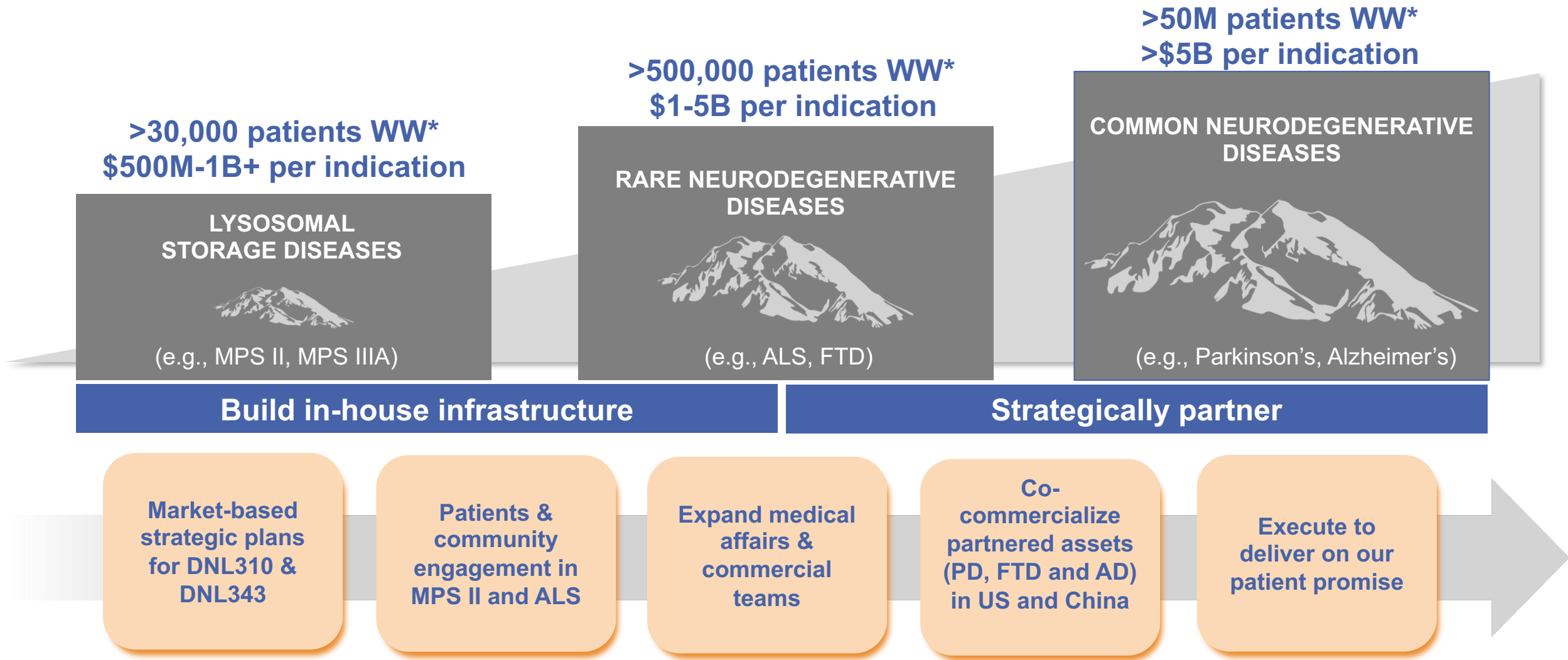
Total earned & potential milestones

~50%

Profit sharing in key geographies

Strategic collaborations facilitate development of a broad portfolio while maintaining commercial upside

OUR VISION: COMMERCIAL ORGANIZATION TO SERVE PATIENTS



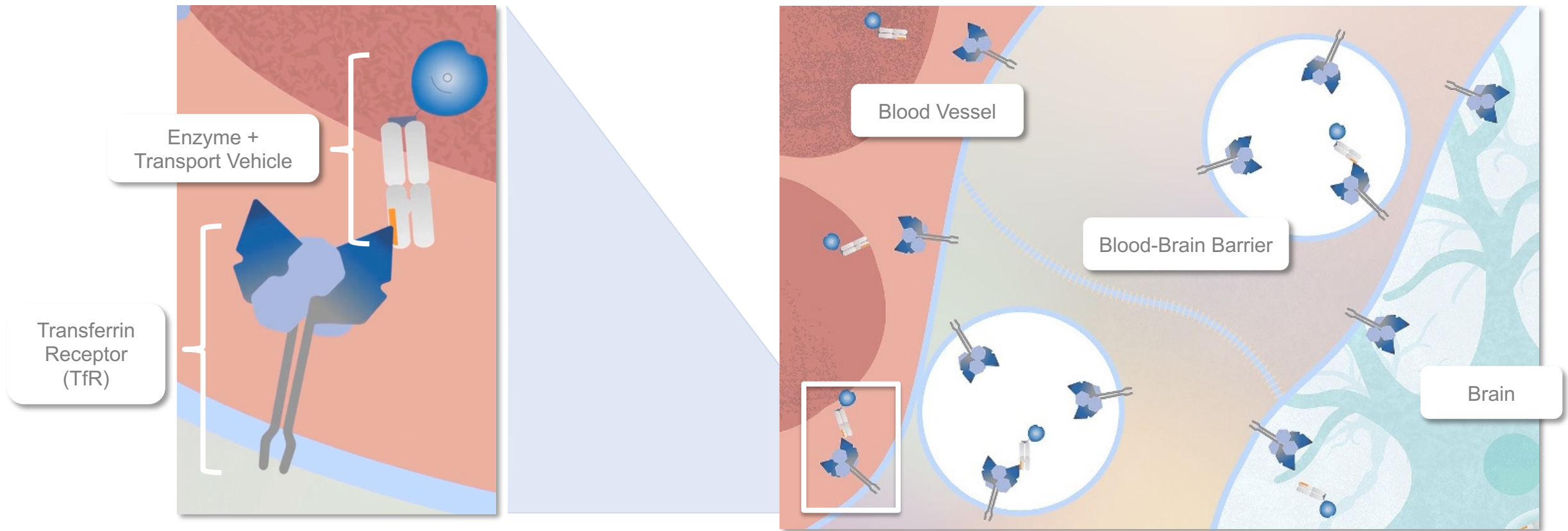
* Denali estimates of world-wide aggregate prevalence



OUR **TV PLATFORM** FOR BRAIN
DELIVERY OF BIOTHERAPEUTICS

ADDRESSING THE CHALLENGE OF DELIVERING THERAPY TO THE BRAIN

The Transport Vehicle (TV) is engineered to deliver efficacious concentrations of biotherapeutics (large molecules) to brain cells via receptor mediated transcytosis

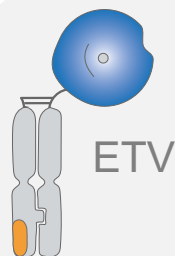


<https://www.denalitherapeutics.com/patients>

TRANSPORT VEHICLE ENABLES MODALITY-OPTIMIZED BRAIN DELIVERY

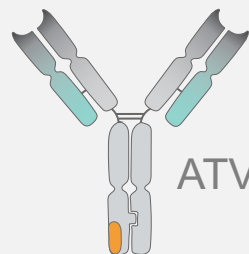
Enzyme Transport Vehicle

Deliver **enzymes** to the brain to replace deficient or missing enzyme activity



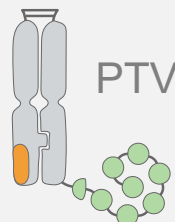
Antibody Transport Vehicle

Deliver **antibodies** in bivalent or bispecific format to the brain



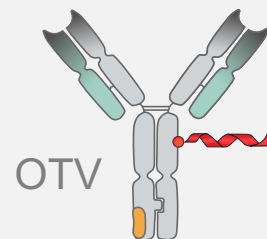
Protein Transport Vehicle

Deliver **proteins** to the brain to replace deficient or missing protein



Oligonucleotide Transport Vehicle

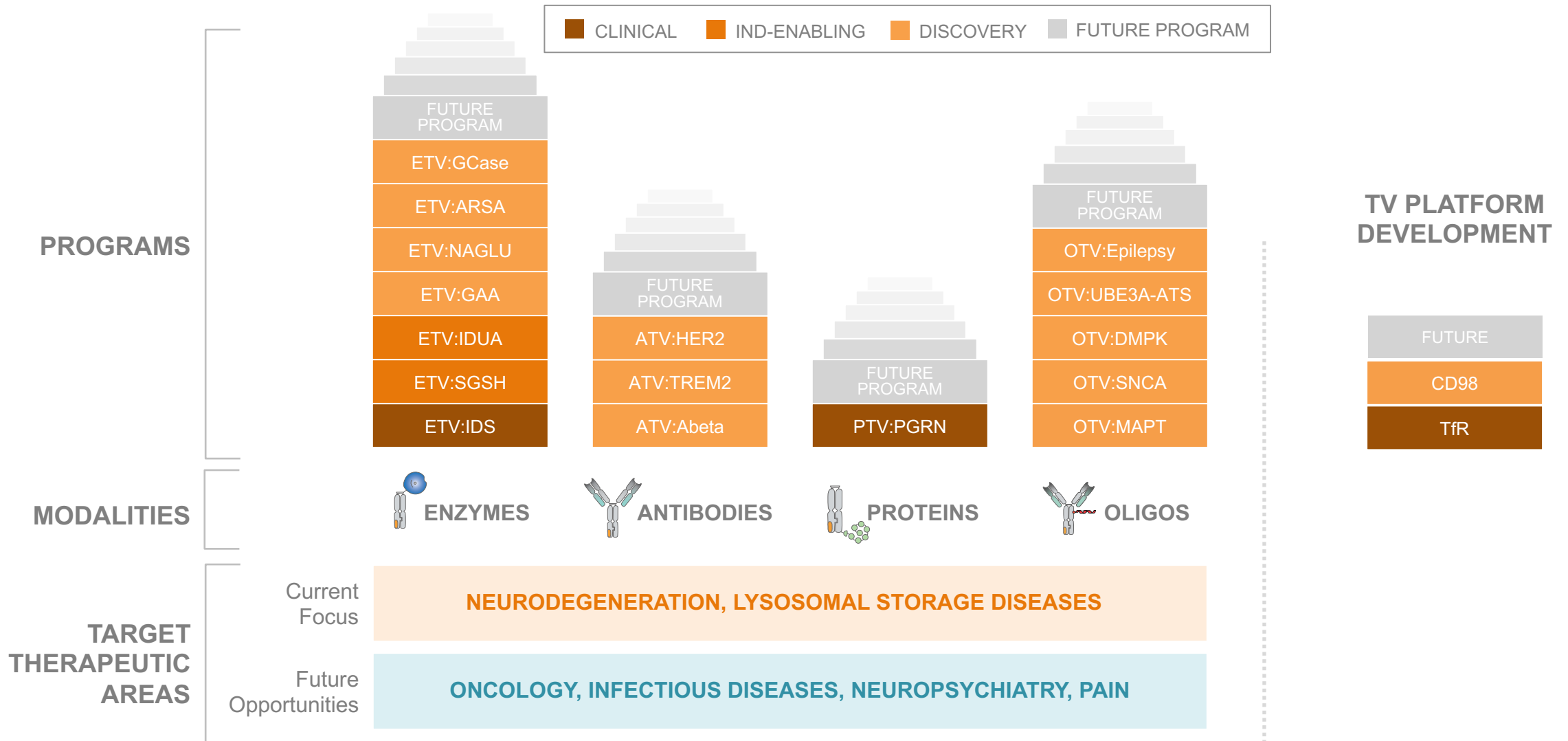
Deliver **oligonucleotides** to the brain and modify gene expression



Each TV modality is a platform opportunity

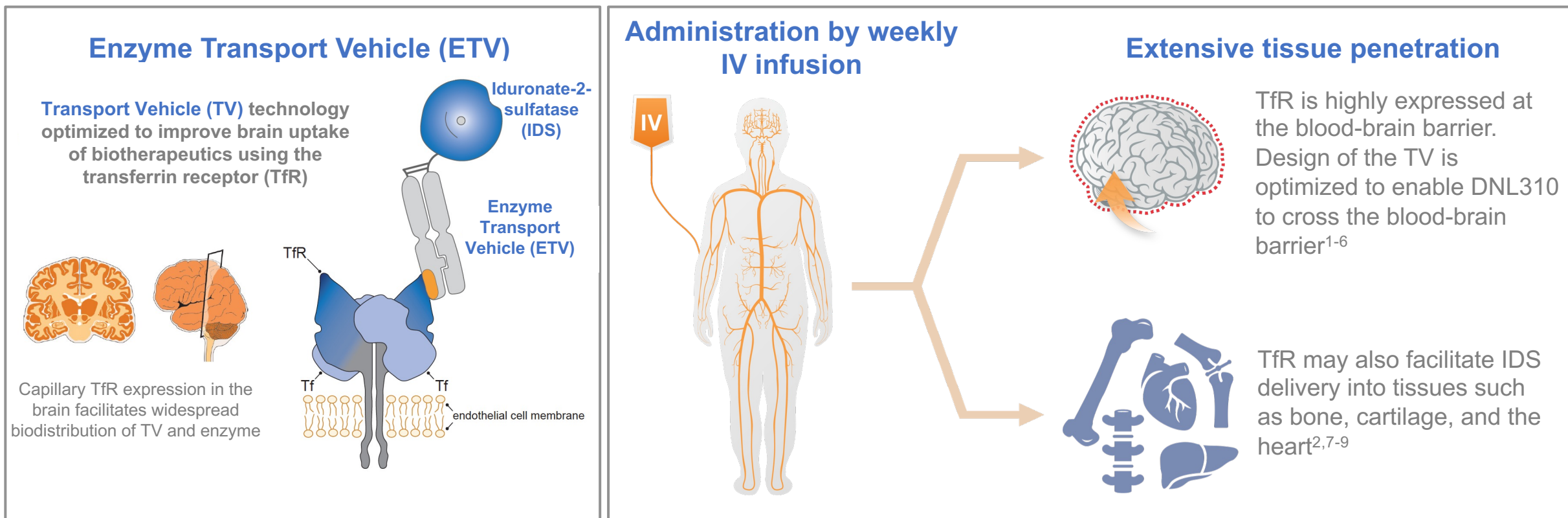
TV PLATFORM OPPORTUNITIES DRIVE SUSTAINABLE VALUE CREATION

Each TV modality is a platform opportunity



DEVELOPING A THERAPY FOR MPS II (HUNTER SYNDROME)

DNL310 (ETV:IDS) is an investigational iduronate-2-sulfatase (IDS) fusion protein engineered to treat both the brain and physical manifestations of mucopolysaccharidosis type II (MPS II) with a **single weekly IV infusion**



DNL310 (ETV:IDS) has the potential to treat neuronopathic and physical manifestations of MPS II

IV, intravenous.

1. Jefferies WA, et al. *Nature*. 1984. 2. Qian ZM, et al. *Pharmacol Rev*. 2002. 3. Bakardjiev AI, et al. *Mol Genet Metab*. 2021. 4. Arguello A et al. *JCI Insight*. 2021. 5. Arguello A, et al. *J Exp Med* 2022. 6. Ullman JC, et al. *Sci Transl Med*. 2020. 7. Wang S, et al. *Haematologica*. 2020. 8. Gammella E, et al. *Metallomics*. 2017. 9. Carlevaro MF, et al. *J Cell Biol*. 1997.

CLINICAL PHENOTYPE OF MPS AND GAG ACCUMULATION

TYPE	NAME	ENZYME DEFICIENCY	GAG
MPS I	Hurler / Scheie	α -L-iduronidase	HS, DS
MPS II	Hunter	Iduronate-2-sulfatase	HS, DS
MPS IIIA	Sanfilippo A	Heparan sulfamidase	HS
MPS IIIB	Sanfilippo B	N-acetyl- α -D-glucosaminidase	HS
MPS IIIC	Sanfilippo C	Acetyl-CoA: α -glucosaminidase	HS
MPS IIID	Sanfilippo D	N-acetylglucosamine-6-sulfatase	HS
MPS IVA	Morquio A	N-acetylgalactosamine-6-sulfatase	KS, CS
MPS VI	Maroteaux-Lamy	N-acetylgalactosamine-4-sulfatase	DS, CS
MPS VII	Sly	β -Glucuronidase	HS, DS, CS
MPS IX	Natowicz	Hyaluronidase	HA

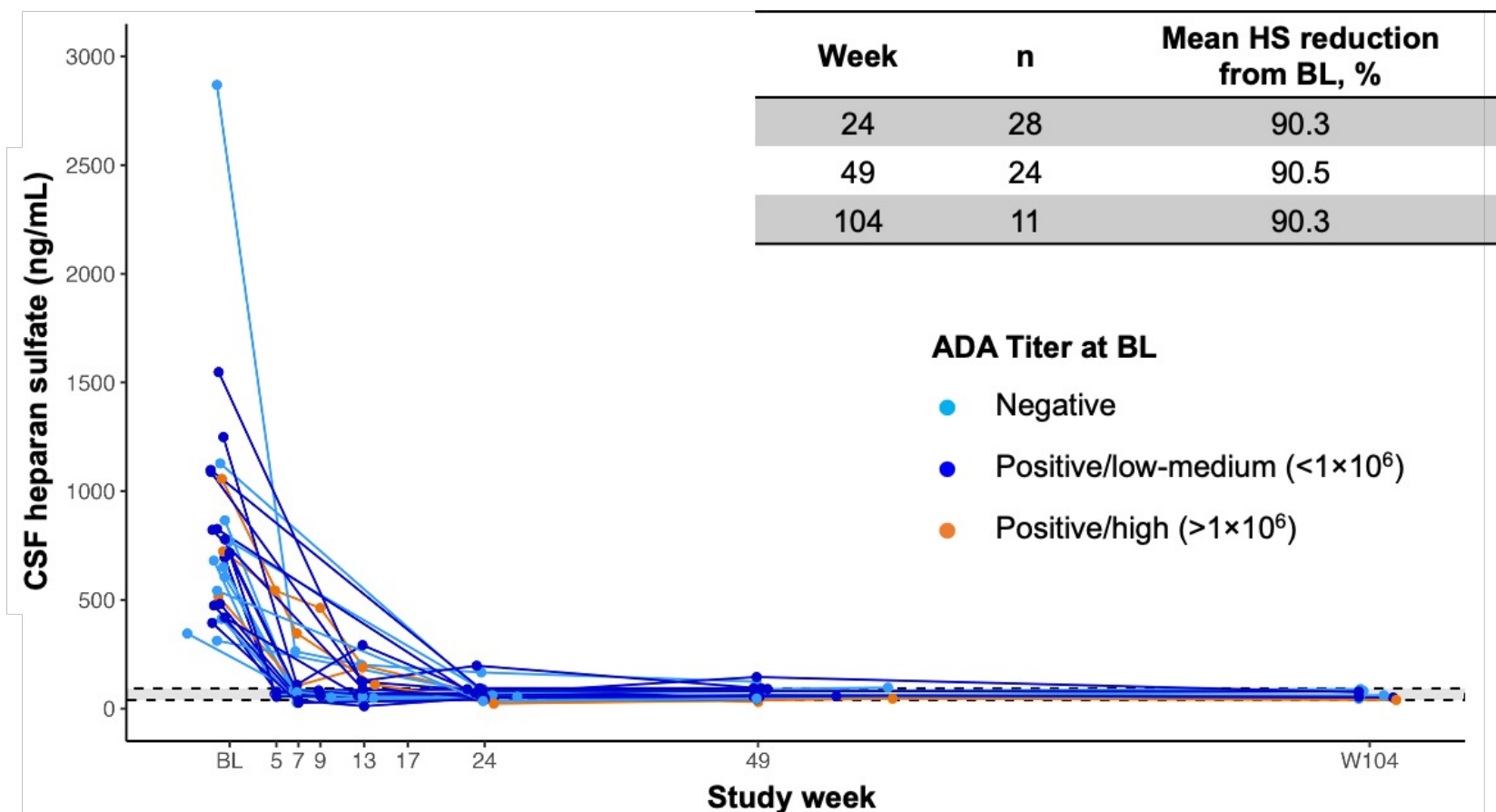
 **CNS involvement**

GAG= glycosaminoglycan
 HS= heparan sulfate
 DS= dermatan sulfate
 CS= chondroitin sulfate
 KS= keratin sulfate
 HA= hyaluronic acid

Heparan sulfate is associated with MPS disorders with CNS involvement

Kobayashi et al., Journal of Human Genetics 2019

DNL310 PHASE 1/2 STUDY BIOMARKERS: CSF HS

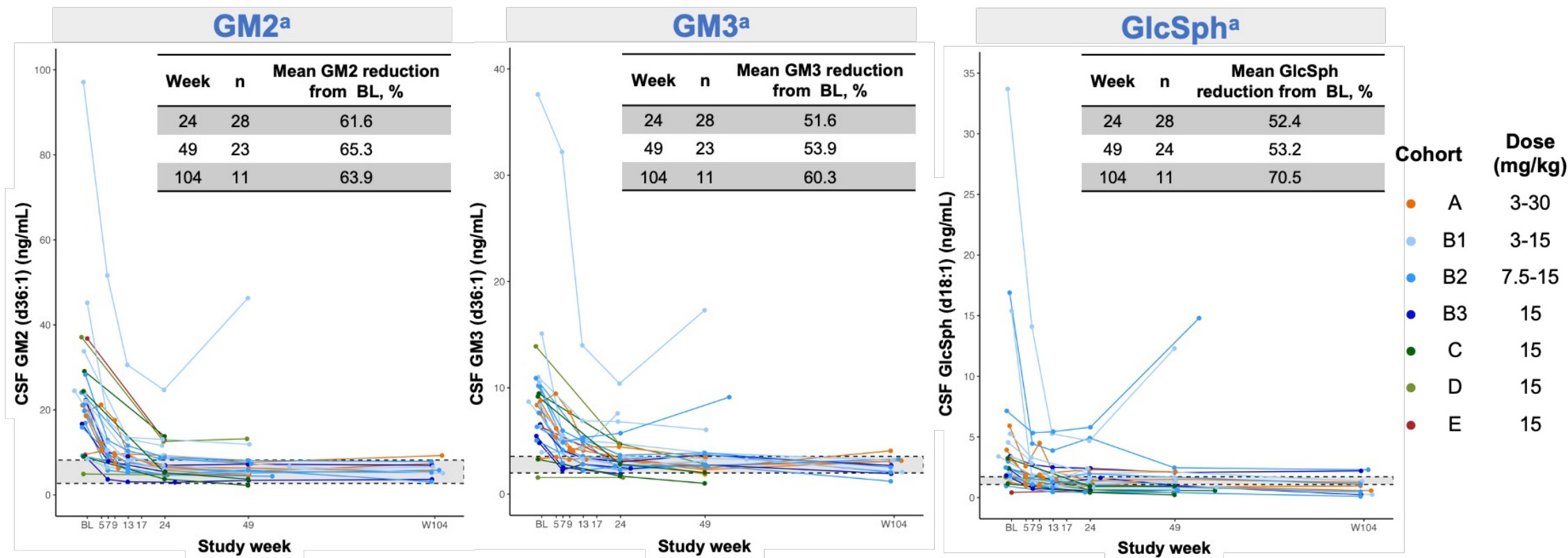


Achievement of normal levels of CSF HS,^a sustained over time, including in those with high pre-existing ADA

ADA, anti-drug antibody; BL, baseline; CSF, cerebral spinal fluid; HS, heparan sulfate.

^aPreliminary normal range (10th and 90th percentile) determined using 30 healthy adult CSF samples (age range, 18-81 years; median, 52 years). Total CSF GAG levels were similar in adults and children (Hendriksz et al. 2015). Normal range for CSF HS, 39.1-92.51 ng/mL. HS was measured as a sum of the disaccharides D0A0, D0A6, D0S0, D2S6.

DNL310 PHASE 1/2 STUDY BIOMARKERS: CSF LYSOSOMAL LIPIDS



Sustained normal levels of CSF lysosomal lipids in most participants are consistent with improved lysosomal function

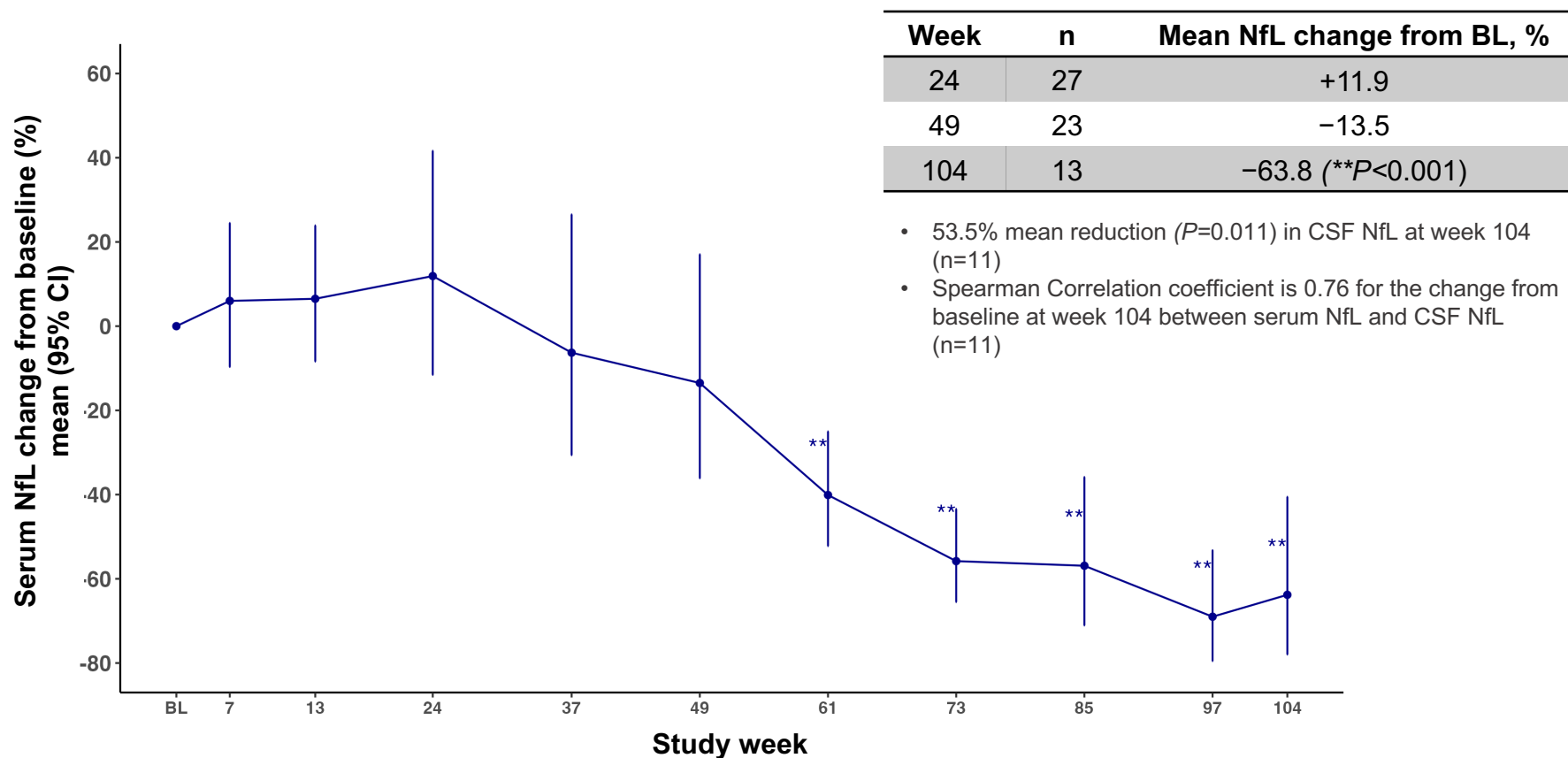
BL, baseline; CSF, cerebral spinal fluid; GlcSph, glucosylsphingosine; GM, ganglioside.

^aPreliminary GM3 normal range (10th and 90th %ile gray dashed lines) determined using 17 healthy adult CSF samples (age range 22-50 years, median 27 years; ng/mL): 1.99-3.55;

Preliminary normal range (10th and 90th %ile gray dashed lines) determined using 18 healthy adult CSF samples (age range 19-52 years, median 24.5 years); GM2 (ng/mL): 2.72-8.2.

GlcSph (ng/mL): 1.08-1.72.

DNL310 PHASE 1/2 STUDY BIOMARKERS: SERUM NEUROFILAMENT (NFL)



Robust reduction in serum NfL, a marker of neuronal damage, significant after 61 weeks and reaching a 64% reduction after two years of dosing with DNL310

** $P<0.001$.

BL, baseline; CSF, cerebral spinal fluid; NfL, neurofilament light chain.

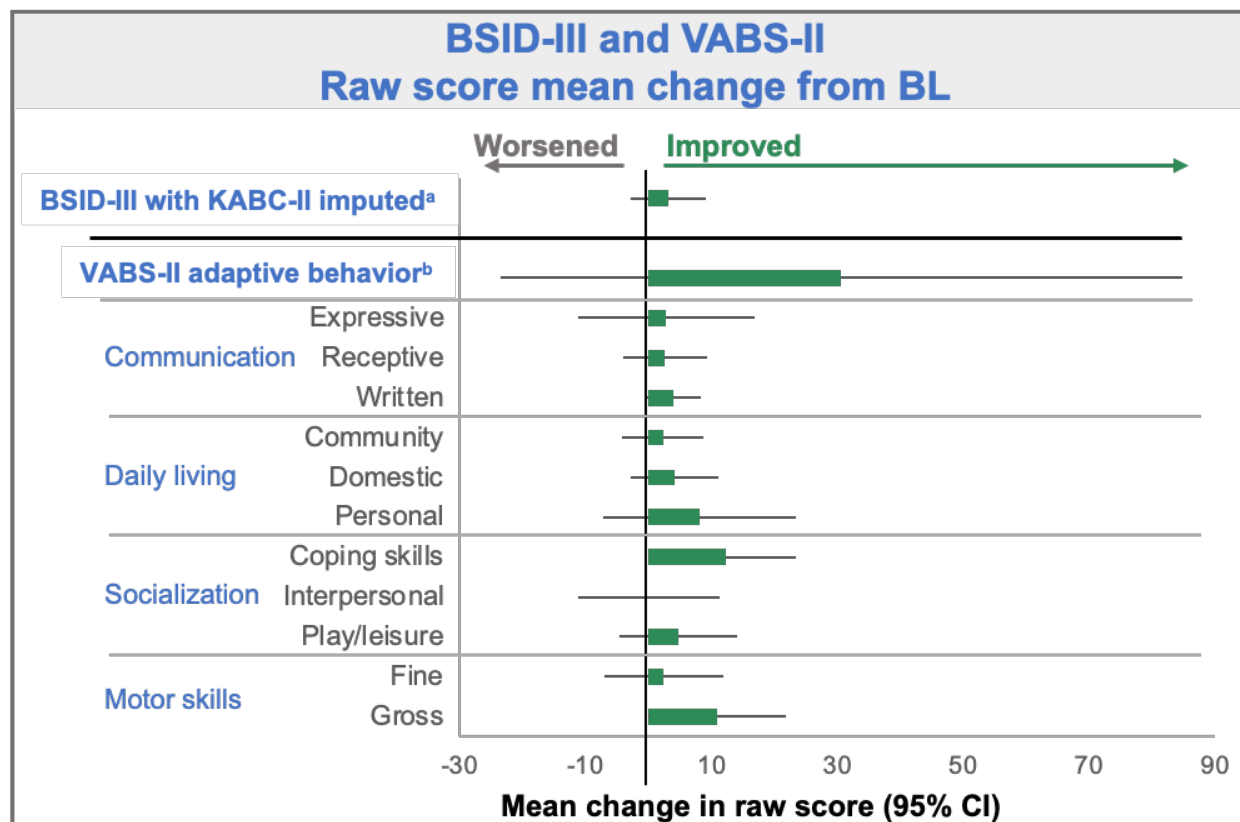
Aggregate summaries by time point are provided for analysis visits that are common across all cohorts. The Week 7 analysis visit includes observations closest to the target day (i.e. Day 43) from weeks 5, 7, or 9. Mean change from baseline are computed from the geometric mean ratio relative to baseline. Corresponding 95% CI and P values are derived from the log ratio relative to baseline. Percent change from baseline are derived as $100(\exp(x)-1)$; where x denotes the mean ratio, upper and lower limit for the mean ratio.

NEUROFILAMENT (NfL): A MARKER OF NEUROAXONAL DAMAGE

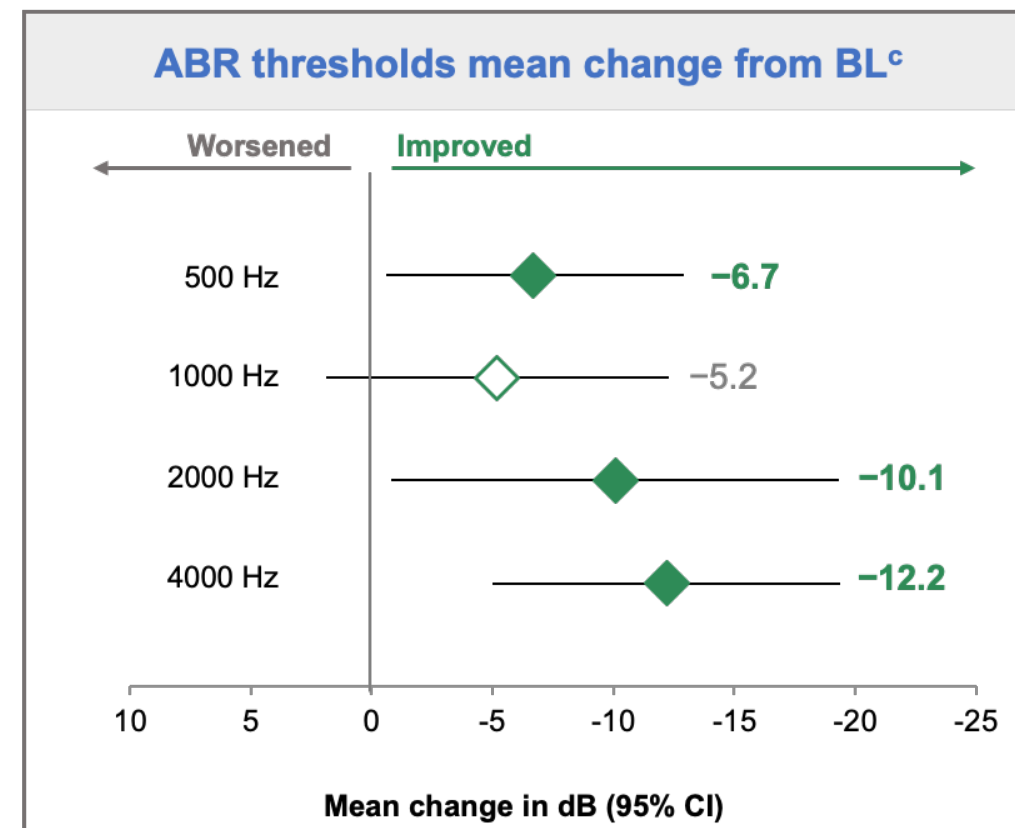
Indication	NfL elevation disease vs. non-disease control	Therapeutic	NfL reduction on treatment	FDA approval
CLN2 ^a	~50-fold (plasma)	cerliponase alfa	~85% (plasma @ 3 yrs)	✓
SMA Type 1 ^b	~30-fold (CSF)	nusinersen	75% (CSF @ ~ Wk 12)	✓
SOD1 ALS ^{c,d}	~4-fold (serum)	tofersen	55% (plasma @ Wk 28)	✓ Accelerated approval
RRMS ^{e,f}	~2-3-fold (plasma)	ocrelizumab interferon beta-1a fingolimod	44% (serum @ Wk 96) 31% (serum @ Wk 96) 43% (plasma @ Wk 52)	✓
PPMS ^e	~2-3-fold (plasma)	ocrelizumab	19% (plasma @ Wk 120)	✓
MPS II^g (neuronopathic)	~5-fold (serum)	DNL310 (ETV:IDS)	64% (serum @ Wk 104)	

- a. Ru Y, et al. "Neurofilament light is a treatment-responsive biomarker in CLN2 disease." *Ann Clin Transl Neurol*. 2019 Dec;6(12):2437-2447.
- b. Olsson B, et al. "NFL is a marker of treatment response in children with SMA treated with nusinersen." *J Neurol* 2019 Sep;266(9):2129-2136.
- c. Halbgebauer, S et al. "Comparison of CSF and serum neurofilament light and heavy chain as differential diagnostic biomarkers for ALS" *Neurodegeneration* 2022; 93, 68-74
- d. Tofersen Prescribing Information
- e. 2020 8TH Joint ACTRIMS-ECTRIMS, Ocrelizumab Treatment Induces a Sustained Blood NfL Reduction in Patients with PPMS and RMS, P0125
- f. Kuhlke, et al. "Blood neurofilament light chain as a biomarker of MS disease activity and treatment response." *Neurology* 2019 Mar 5; 92(10): e1007–e1015
- g. Bhalla A, et al. "Characterization of Fluid Biomarkers Reveals Lysosome Dysfunction and Neurodegeneration in Neuronopathic MPS II Patients." *Int. J. Mol. Sci.* 2020, 21, 5188

DNL310 PHASE 1/2 STUDY: CLINICAL OUTCOMES AT WEEK 49



(n=16)



(n=18 to 20 participants, varies by frequency)

Improvements in mean cognitive BSID-III and VABS-II raw scores, and ABR thresholds at week 49 of DNL310 treatment suggest positive effects on cognition, adaptive behavior, and hearing

ABR, auditory brainstem response; BL, baseline; BSID-III; Bayley Scales of Infant and Toddler Development III; KABC, Kaufman Assessment Battery for Children; VABS-II, Vineland Adaptive Behavior Scales II.

^aImputed values are taking a value of 91. Participants with imputed values at W49 and Baseline are not considered in the mean change as both values are 91 leading to an undesired change values of 0. ^bData from 4 participants either unavailable (n=1) or only VABS-3 collected (n=3) at Week 49. The Total Adaptive Behavior raw score derives from all Communication, Daily Living, and Socialization subdomains except for Communication-Written, Daily Living-Domestic, and Daily Living-Community. ^cResults are based on air conduction tests. Least squares mean (95% CI), adjusted for age at ERT initiation.

Source: Burton B, et al. SSIEM 2023

DNL310 PHASE 1/2 STUDY INTERIM SAFETY: OVERVIEW

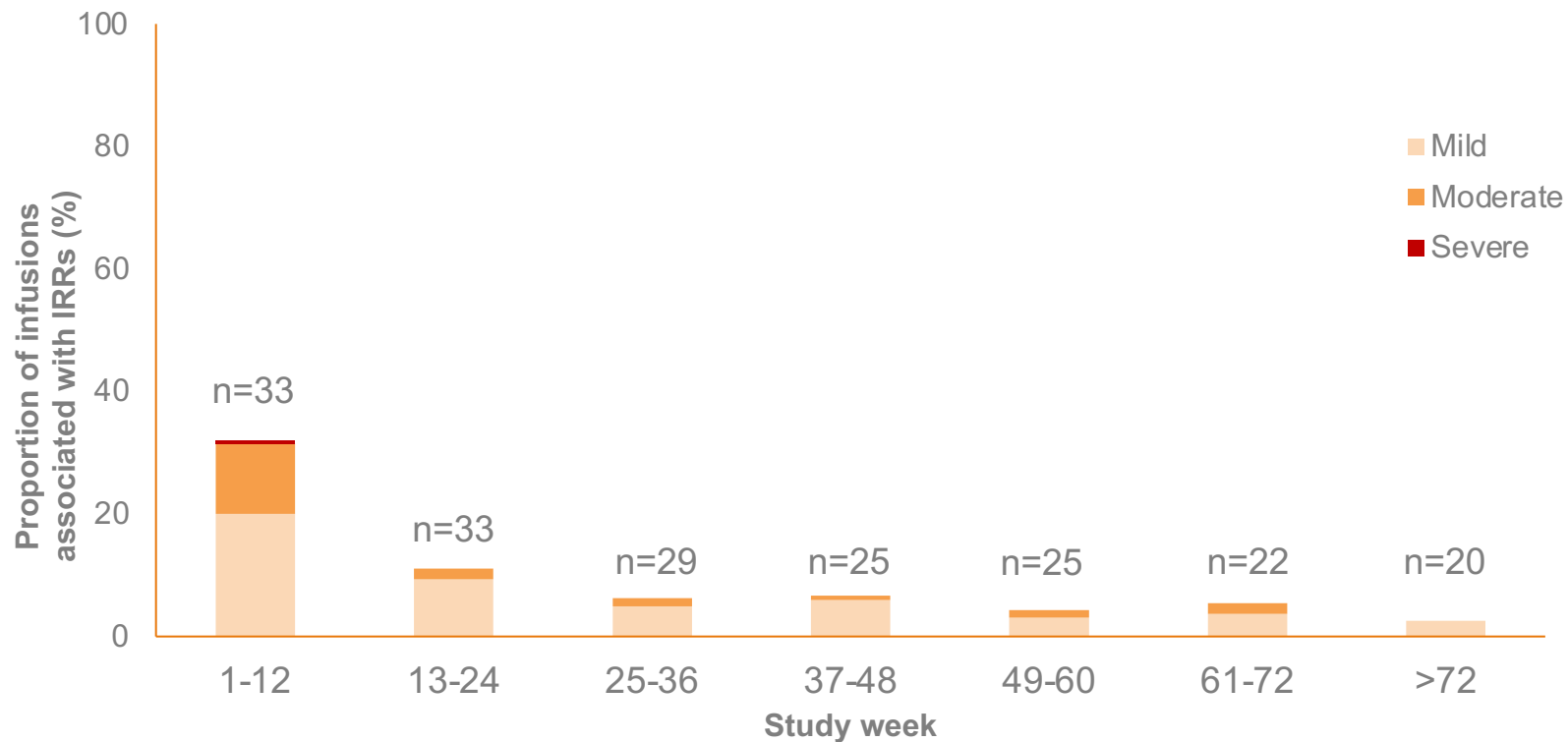
Cumulative information, including previously reported^{1,2}

TEAEs	<ul style="list-style-type: none">• All participants reported treatment-emergent adverse events (TEAEs), which were mostly mild or moderate• There were no dose-related safety findings• Infusion-related reactions (IRRs) were the most frequent TEAEs, IRR frequency declines over time (see next slide)• Adverse events of special interest (AESIs) were as follows:<ul style="list-style-type: none">• 20 participants experienced moderate IRRs, and 1 participant experienced severe IRRs• 4 participants (all with mild baseline anemia or a history of anemia) had moderate anemia (3 resolved); dosing continued in all 4 cases; anemia is a known complication of lysosomal storage diseases such as MPS II³• One discontinuation related to TEAEs (including IRRs and other non–drug-related AEs) was observed in a participant with complex underlying disease; 3 other discontinuations due to social reasons (family circumstances, relocation)
SAEs	<ul style="list-style-type: none">• SAEs were reported in 10 participants; of these, 2 had IRRs, and 8 had SAEs that are largely known comorbidities of MPS II or childhood infections and are unrelated (per the investigators) to study drug or procedures (including constipation, upper respiratory tract infection, progressive cervical stenosis/thoracic syrx, increased episodes of OSA, vomiting and diarrhea, viral parotitis, central line infection)
SAFETY LABS	<ul style="list-style-type: none">• Prior to treatment, 15 participants had elevated total urine GAGs (colorimetric assay); all decreased after receiving DNL310• No other significant trends in safety laboratory evaluations occurred post initiation of DNL310 treatment

Safety profile reflects median treatment duration of 91 weeks in 33 study participants; maximum treatment duration: 135 weeks
Independent Data Monitoring Committee recommended continuing study without modifications (May 2023)

DNL310 is generally well tolerated with a safety profile that continues to support development in MPSII

DNL310 PHASE 1/2 STUDY SAFETY: IRRS



Total number of infusions during the study: 2471

Tolerance to DNL310 occurred with longer-term dosing

DNL310 PHASE 1/2 STUDY: SUMMARY OF INTERIM RESULTS

Clinical safety

- Safety profile is based on 33 participants with MPS II with a median treatment duration of 91 weeks, and supports continued development in MPS II
- IRRs accounted for the most frequent TEAEs and decreased in frequency and severity with continued dosing

Biomarkers

- Rapid normalization or near normalization of CSF HS was observed in all participants, sustained at weeks 49 and 104
- Normalization of CSF HS was observed even in participants with high preexisting ADA
- Normalization of CSF lysosomal lipids in most participants consistent with improved lysosomal function
- Robust reduction of 64% in serum NfL, a marker of neurodegeneration, with long-term dosing

Clinical outcomes

- Interim clinical outcomes data suggest positive change in adaptive behavior and cognition with DNL310 treatment
- ABR data suggest that DNL310 treatment improves auditory function

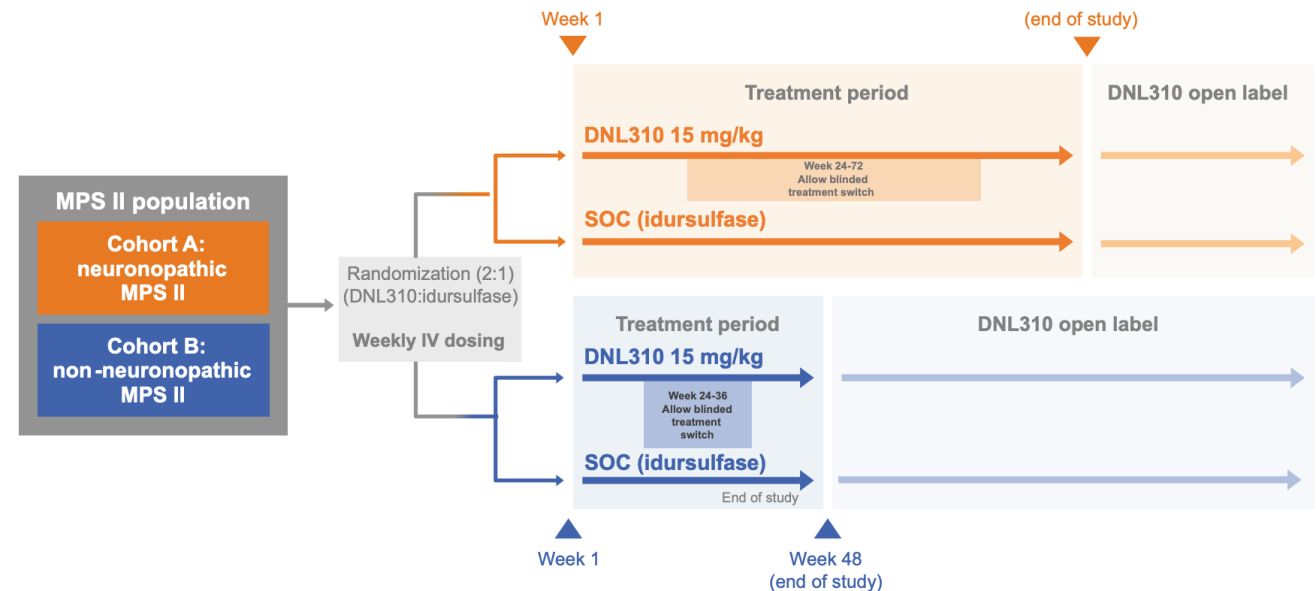
- **DNL310 is an investigational, intravenously administered, Enzyme Transport Vehicle (ETV)-enabled, iduronate-2-sulfatase (IDS) replacement therapy designed to cross the BBB and address the behavioral, cognitive and physical manifestations of MPS II**
- **A potentially registrational Phase 2/3 study with sites in North America, South America, and Europe is enrolling (NCT05371613)**

DNL310 PHASE 2/3 STUDY DESIGN IN PEDIATRIC MPS II PATIENTS

DNLI-E-0007 STUDY OVERVIEW (NCT05371613)

Study Design	Double-Blind, Randomized Study of DNL310 vs Idursulfase in children with neuronopathic (96-week study) or non-neuronopathic (48-week study) MPSII followed by OLE <ul style="list-style-type: none"> DNL310 is administered by weekly IV infusion n= 54 patients in 2 cohorts
Key Eligibility	<ul style="list-style-type: none"> Cohort A (n=33): neuronopathic patients aged ≥2 to <6 years Cohort B (n=21): non-neuronopathic patients aged ≥6 to <17 years Receiving approved IDS for >4 months IDS-treated patients will be switched to DNL310 without a washout period
Key Endpoints	Key Efficacy Endpoints <ul style="list-style-type: none"> Effect of DNL310 on CSF biomarkers <ul style="list-style-type: none"> CSF GAGs Effect of DNL310 on neurobehavioral parameters <ul style="list-style-type: none"> Adaptive behavior testing: Vineland Adaptive Behavior Scales Neurocognitive testing: BSID, KABC, WISC Effect of DNL310 on peripheral manifestations of disease <ul style="list-style-type: none"> Urine GAGs Liver/spleen volume Clinician and caregiver reported outcomes: Global Impression Scales Key Safety Assessments <ul style="list-style-type: none"> Treatment-emergent adverse events Infusion-related reactions Laboratory abnormalities

DOSING SCHEMA



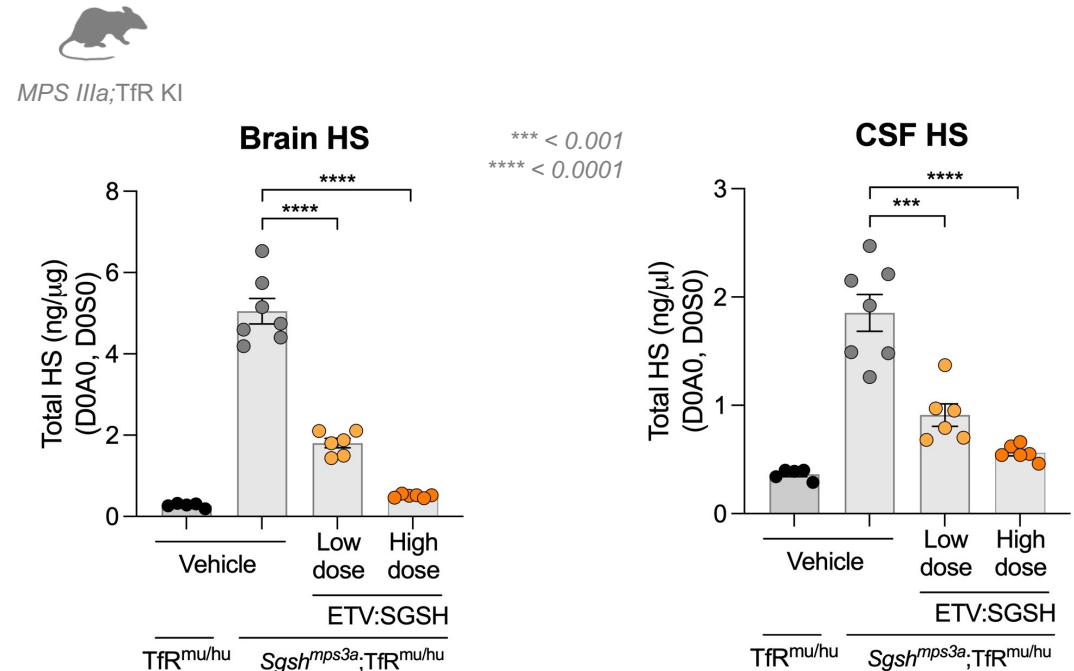
DNL126 (ETV:SGSH): EXPANDS ETV PLATFORM FOR MPS IIIA

Addressing cognitive, behavioral & physical manifestations of Sanfilippo syndrome Type A

- Rare lysosomal storage disease (LSD) that causes neurodegeneration; no treatments
- Caused by genetic mutations that result in a reduction in the activity of SGSH
- SGSH is an enzyme responsible for degrading heparan sulfates (HS) in the lysosome
- HS accumulation leads to lysosomal dysfunction
- DNL126 is designed to replace SGSH in the brain and throughout the body

ETV:SGSH=Enzyme Transport Vehicle N-Sulfoglucosamine Sulfohydrolase;
MPS=mucopolysaccharidoses; CSF=cerebrospinal fluid; IND=investigational new drug

IV DNL126 treatment reduces HS in a dose-dependent manner in brain and CSF



Recruiting activities for the Phase 1/2 study beginning in 2H23

DNL593 (PTV:PGRN): PGRN BRAIN DELIVERY FOR FTD-GRN

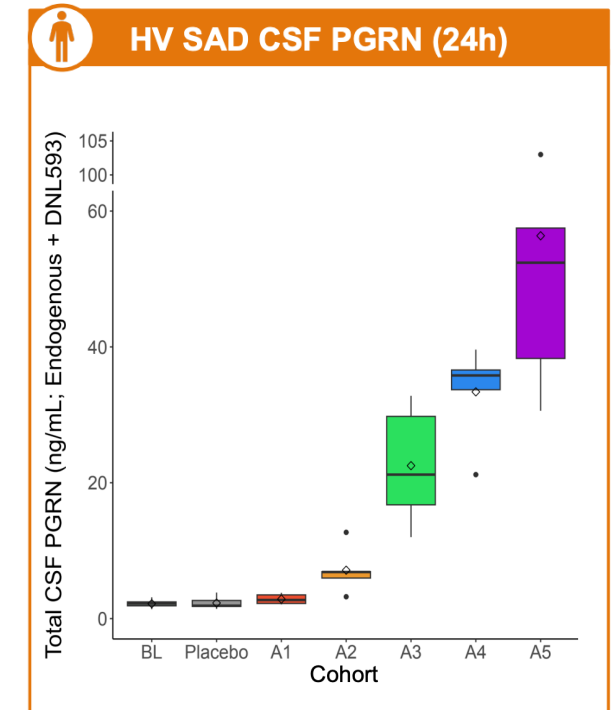
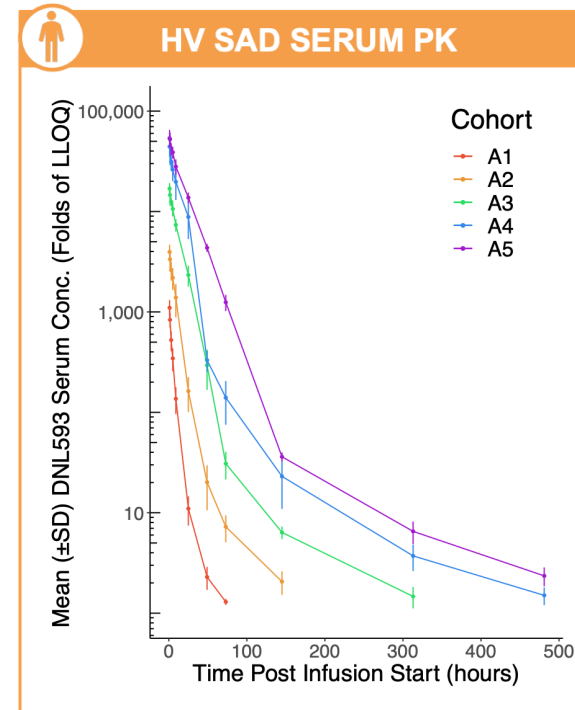
Brain delivery of progranulin (PGRN) designed to treat FTD-GRN

- FTD is the most common dementia in people under 60; no approved therapies
- FTD-GRN is associated with PGRN deficiency; accounts for 5-10% of FTD
- Single doses of DNL593 in HVs led to dose-dependent increases in CSF PGRN and were generally well tolerated
- Data support enrolling participants with FTD-GRN in Part B (multiple ascending doses)
- Co-development/co-commercialization with Takeda



PTV:PGRN=Protein Transport Vehicle:Progranulin; FTD-GRN-frontotemporal dementia granulin; CSF=cerebrospinal fluid; HVs=healthy volunteers

Dose-dependent increase in CSF PGRN in HV with IV DNL593 further validates TV for BBB crossing

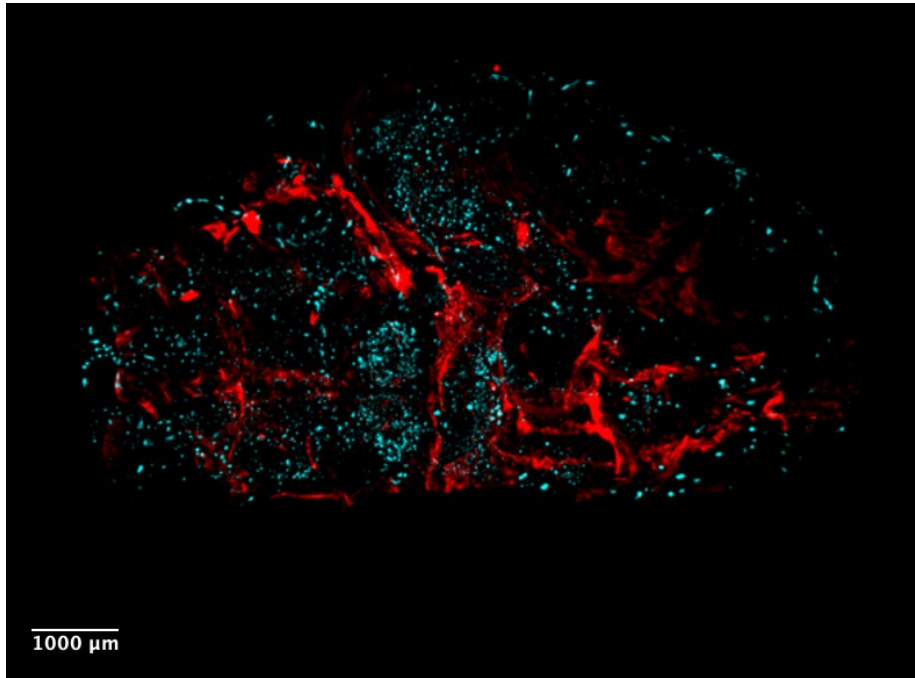


Recruitment of participants with FTD-GRN in Part B (ascending multiple doses) of the Phase 1/2 study is ongoing

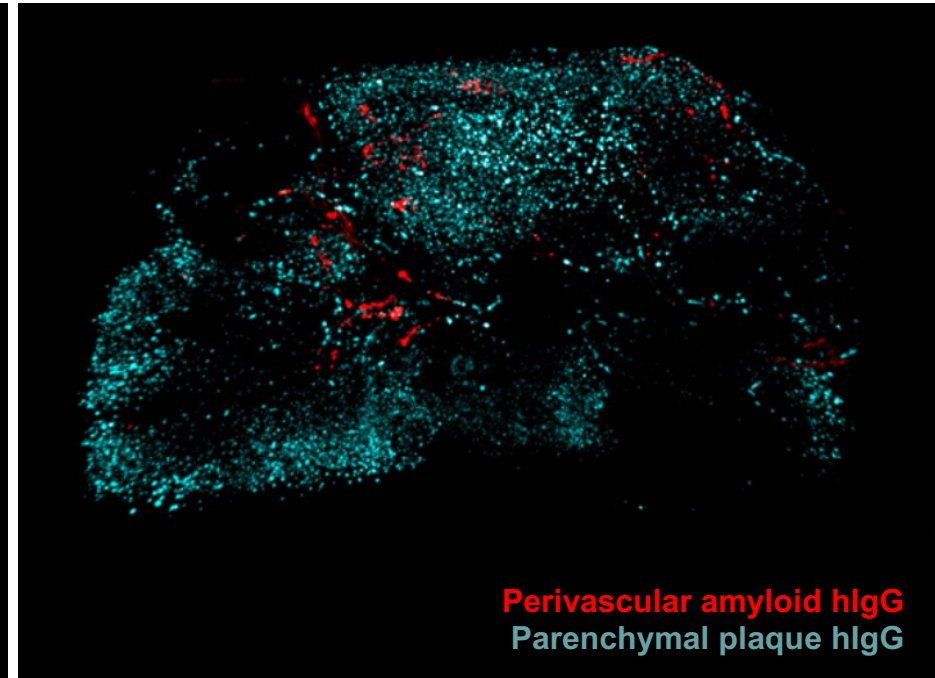
ATV:ABETA FOR ALZHEIMER'S DISEASE (AD)

ATV:Abeta shows broad parenchymal plaque binding with minimal perivascular distribution

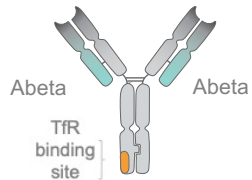
Anti-Abeta



ATV:Abeta



5xFAD; Tfr KI



iDISCO whole brain image 24h post 10mg/kg
single dose in AD mouse model

Biogen has opted-in to the ATV:Abeta program (April 2023) and now leads development and commercialization

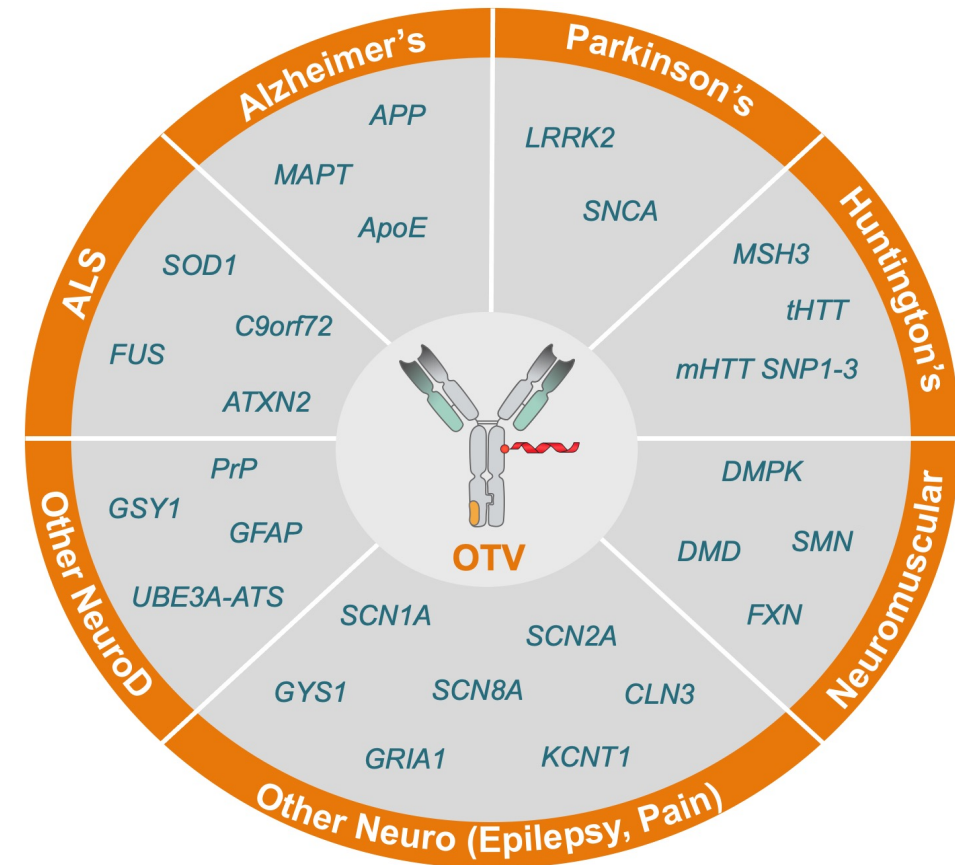
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OTV IS DESIGNED TO ENHANCE CNS DELIVERY OF OLIGONUCLEOTIDES

Therapeutic oligonucleotides have the potential to address challenging targets

- Oligonucleotide Transport Vehicle (OTV) is designed to:
 - Enable superior biodistribution of ASOs across brain regions
 - Provide superior knockdown of target gene expression across all cell types
 - Enable IV dosing
- OTV opens a large potential indication space in neurodegeneration and beyond
- Multiple OTV programs progressing toward IND-enabling studies
- OTV manuscript posted on bioRxiv April 28, 2023 (Barker SJ et al.)

OTV has potential to revolutionize ASOs/oligos for treating CNS disease



Illustrative

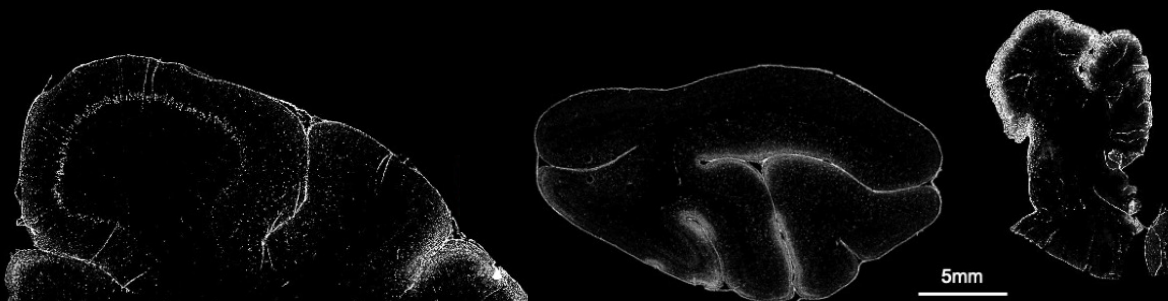
OTV PROVIDES UNIFORM ASO DEPOSITION ACROSS THE CNS WITH IV DELIVERY



NAKED ASO INTRATHECAL (IT) DELIVERY

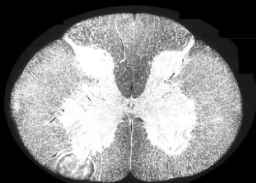
Limited ASO Biodistribution

BRAIN
ANTERIOR → POSTERIOR

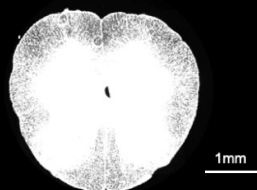


SPINAL CORD

Cervical



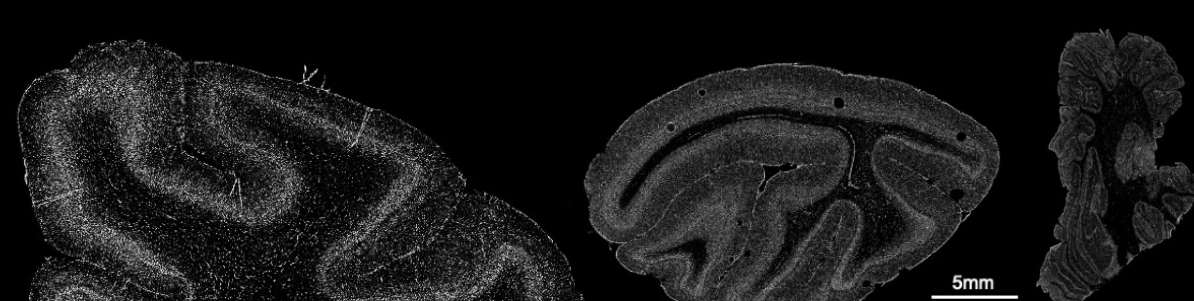
Lumbar



OTV INTRAVENTROUS (IV) DELIVERY

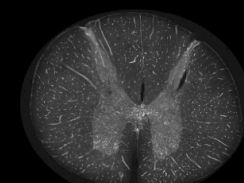
Widespread ASO Biodistribution

BRAIN
ANTERIOR → POSTERIOR

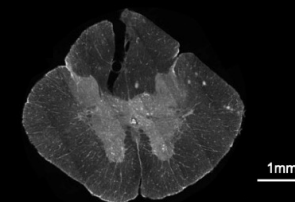


SPINAL CORD

Cervical



Lumbar



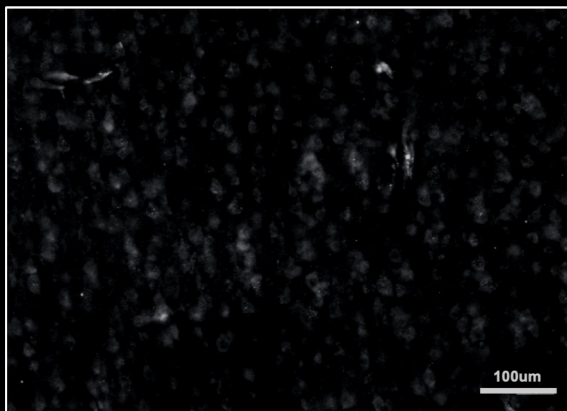
OTV PROVIDES UNIFORM ASO DEPOSITION ACROSS THE CNS WITH IV DELIVERY



NAKED ASO INTRATHECAL (IT) DELIVERY

Limited ASO Biodistribution

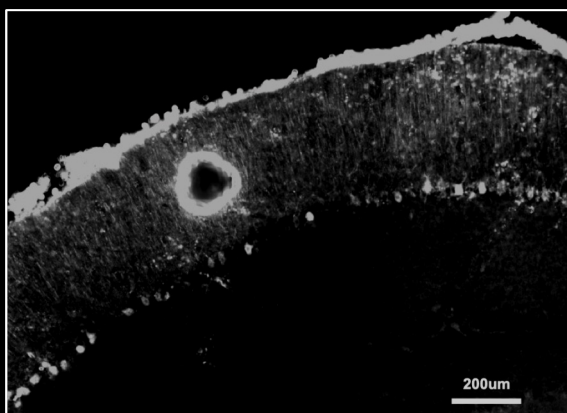
CORTEX



STRIATUM



CEREBELLUM



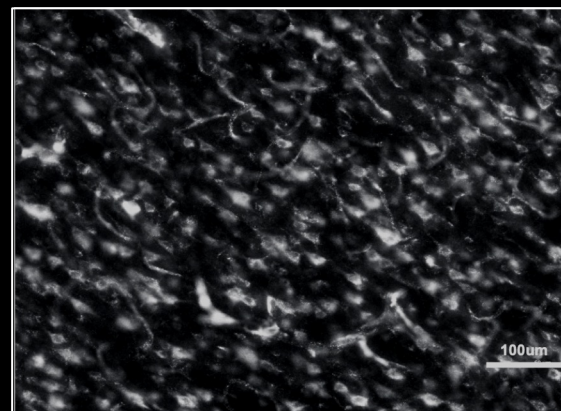
WHITE MATTER



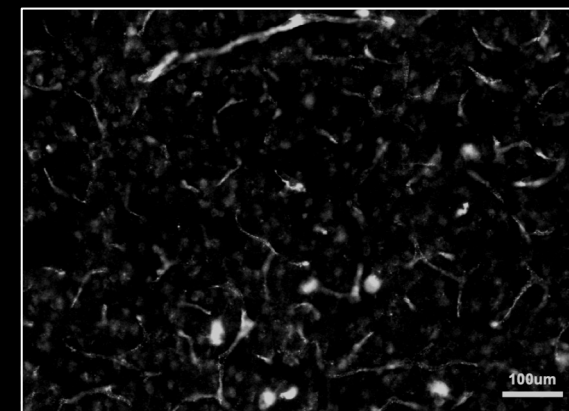
OTV INTRAVENTROUS (IV) DELIVERY

Widespread ASO Biodistribution

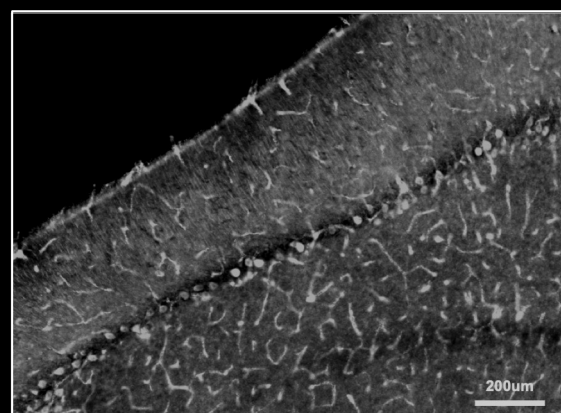
CORTEX



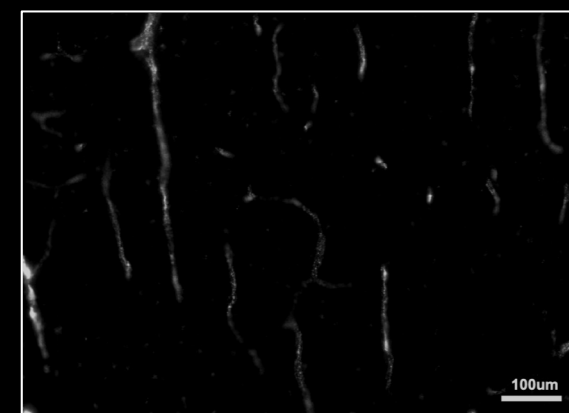
STRIATUM



CEREBELLUM

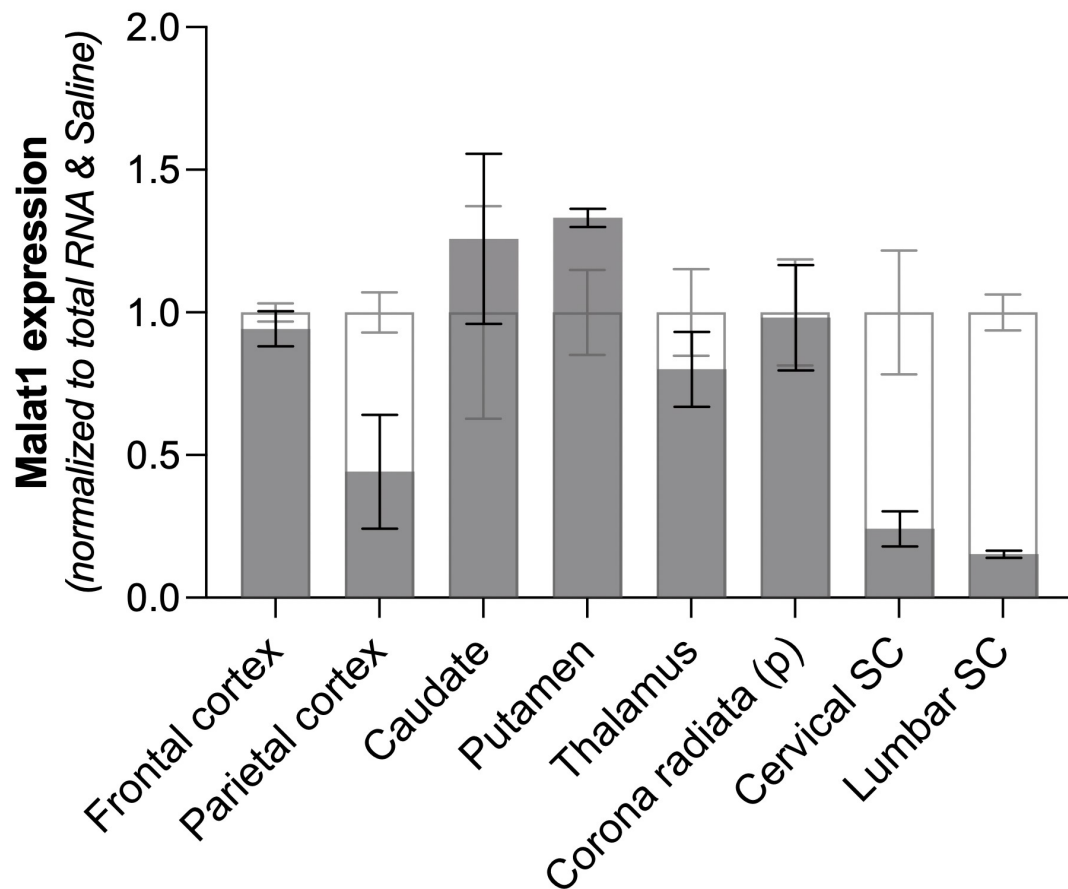


WHITE MATTER

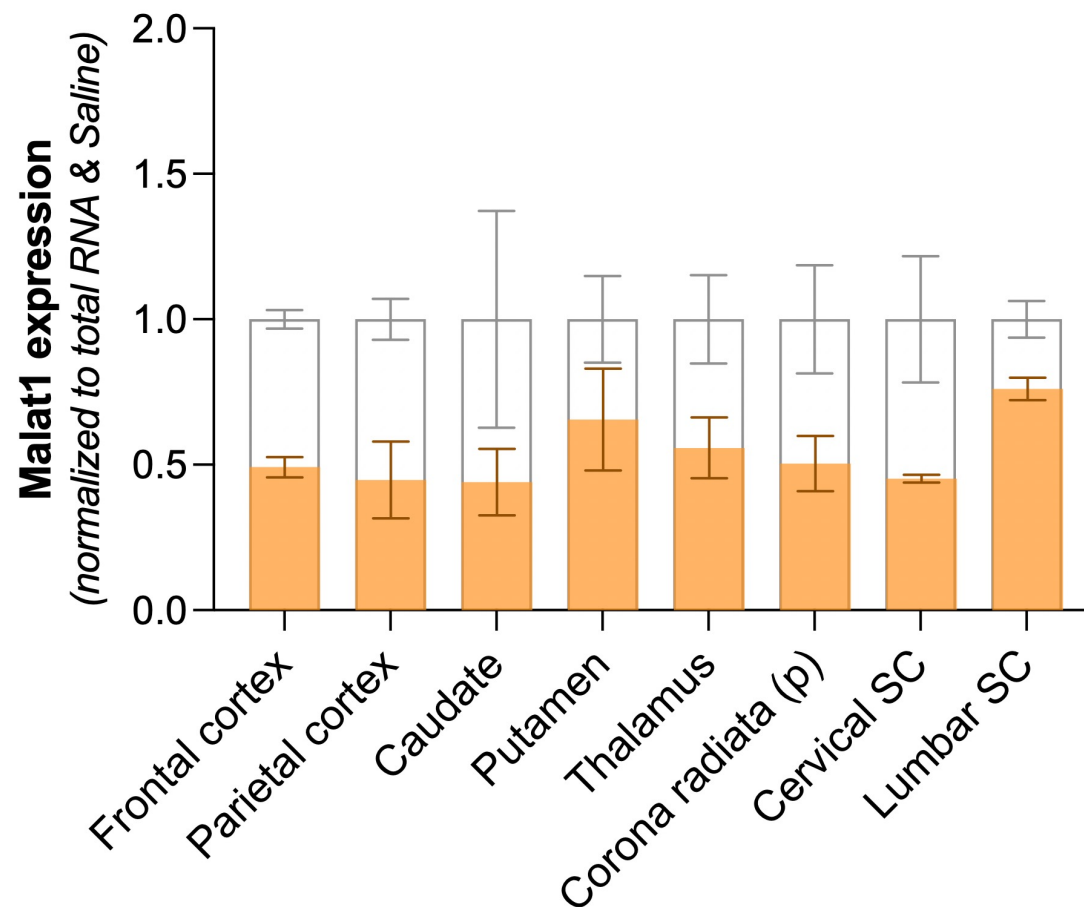


OTV ENABLES MORE UNIFORM KNOCKDOWN OF TARGET GENE EXPRESSION

NAKED ASO INTRATHECAL (IT) DELIVERY



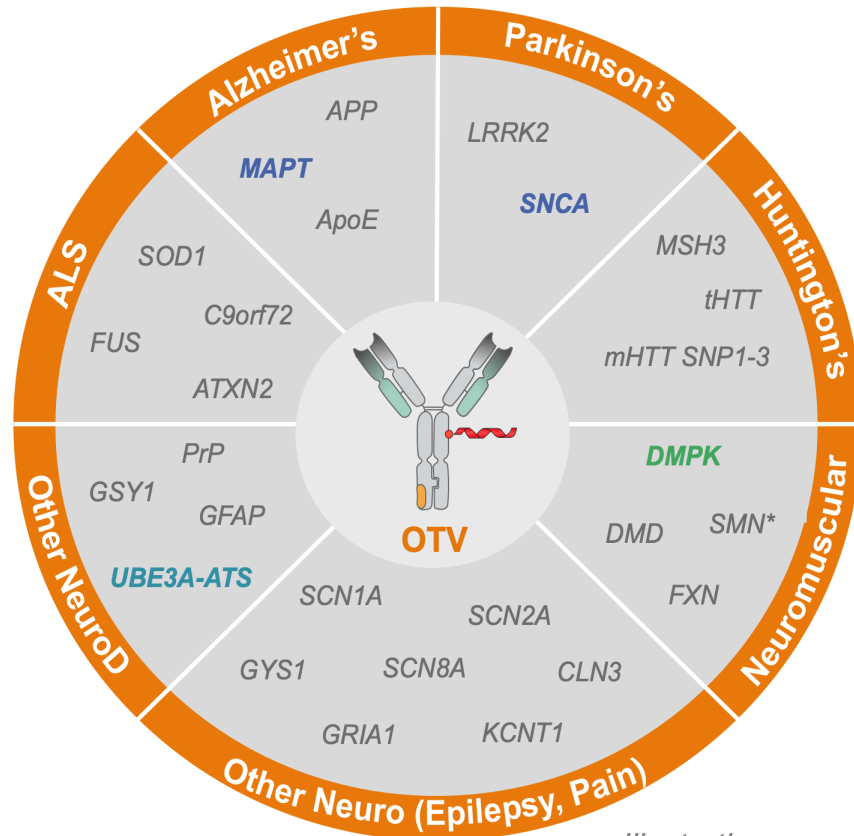
OTV INTRAVENOUS (IV) DELIVERY



Data represented as mean \pm SEM; n=3

IV OTV shows uniform knockdown across the CNS compared to IT ASO

OTV TARGET SELECTION



Illustrative

OTV in IND-Enabling stage with near-term focus on acceleration of two targets to clinical testing

TARGET	INDICATION	PREVALENCE	DIFFERENTIATION STRATEGY
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COMMON NEURODEGENERATIVE DISEASES

MAPT	Alzheimer's Disease	6-10M cases US	Uniform knockdown of MAPT across the CNS to effectively reduce all forms of Tau protein & decrease aggregates
SNCA	Parkinson's Disease	1M cases US	Uniform knockdown of SNCA across the CNS to effectively reduce all forms of α-Syn protein & decrease aggregates

RARE CNS DISEASES

UBE3A-ATS	Angelman's Syndrome	1.5-3K cases US (<8yo)	Uniform knockdown of UBE3A-ATS via systemic route to increase normal UBE3A protein levels throughout the CNS
Epilepsy Target 1	Epilepsy	1-15K cases US	Undisclosed

NEUROMUSCULAR DISEASES

DMPK	Myotonic Dystrophy Type 1	<i>Adult</i> 40K cases US <i>Congenital</i> ~600 cases US	Knockdown of DMPK in periphery and CNS to reduce toxic RNA foci & allow MBNL proteins to resume normal splicing
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OUR BRAIN-PENETRANT **SMALL MOLECULE** PROGRAMS

BIIB122 (LRRK2 INHIBITOR): TARGETING THE LYSOSOME IN PD

Targeting LRRK2 may impact the underlying biology and slow the progression of PD

- 10M+ people with Parkinson's disease (PD) WW
- Mutations in LRRK2 are one of the most common genetic risk factors for PD
- Increased LRRK2 kinase activity is thought to impair lysosomal function and contribute to PD
- Denali conducted extensive Phase 1/1b testing with LRRK2 inhibitors in 300+ individuals*
- BIIB122 achieved $\geq 80\%$ pS935 inhibition (target engagement biomarker) at doses of ≥ 225 mg
- Biogen is leading operational execution of the Phase 2b LUMA Study

Phase 2b LUMA Study of BIIB122 in PD patients with and without LRRK2 mutations

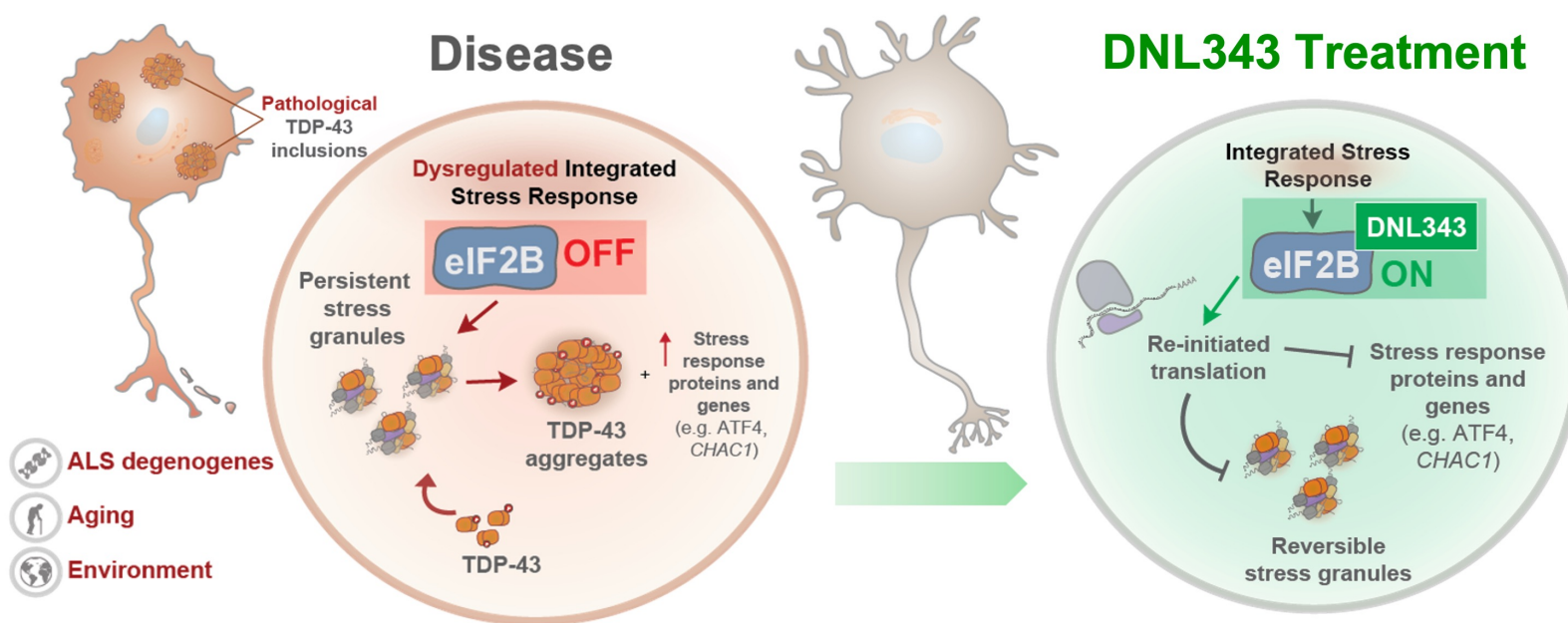
	Phase 2b LUMA Study
PD patient pop.	Early-stage, idiopathic and pathogenic LRRK2 variants
Dosing	225 mg oral once daily BIIB122 vs. placebo
Primary endpoint	Assessed using MDS-UPDRS
No. participants	640 (320 per arm)
Treatment period	48 weeks (min)
Study initiation	May 2022

*Phase 1/1b program for BIIB122 and DNL201

LRRK2=leucine-rich repeat kinase 2; WW=worldwide; MDS-UPDRS=Movement Disorder Society Sponsored Revision of the Unified Parkinson's Disease Rating Scale

EIF2B ACTIVATION HAS POTENTIAL TO SLOW NEURODEGENERATION IN ALS

In ALS, TDP-43 pathology is linked to cellular dyshomeostasis resulting from chronic activation of the Integrated Stress Response (ISR) via inactivation of the eukaryotic initiation factor 2b (eIF2B)

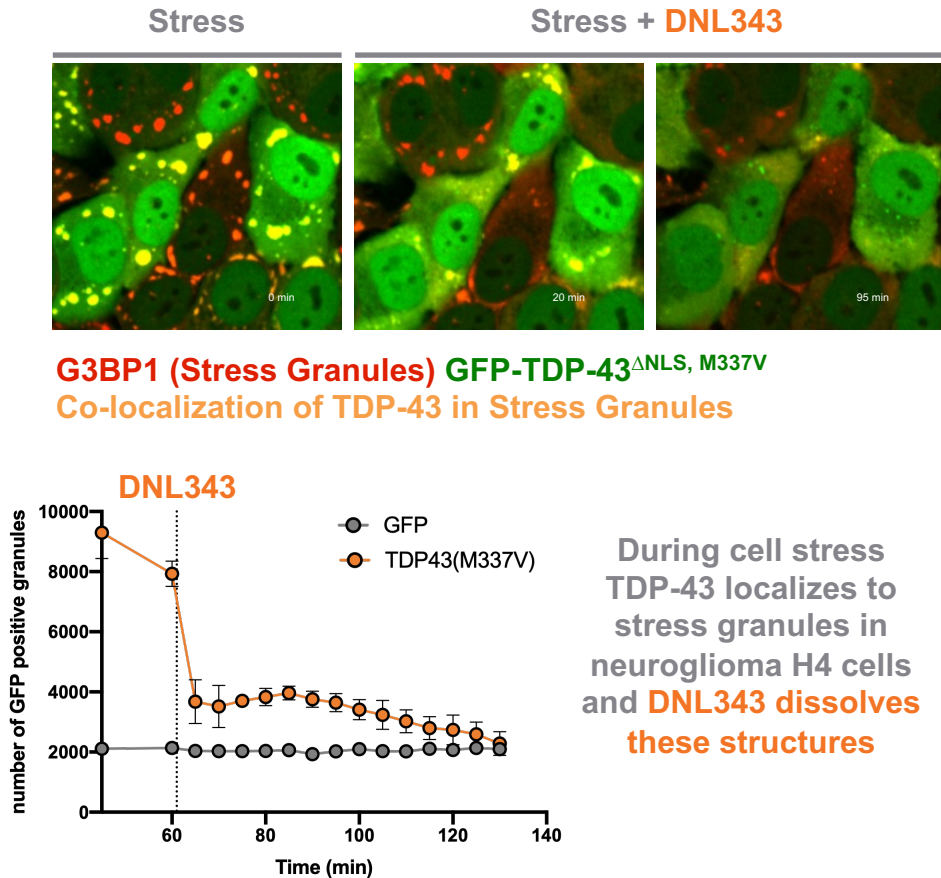


DNL343 is an eIF2B agonist designed to inhibit the ISR and restore cells to a healthy state

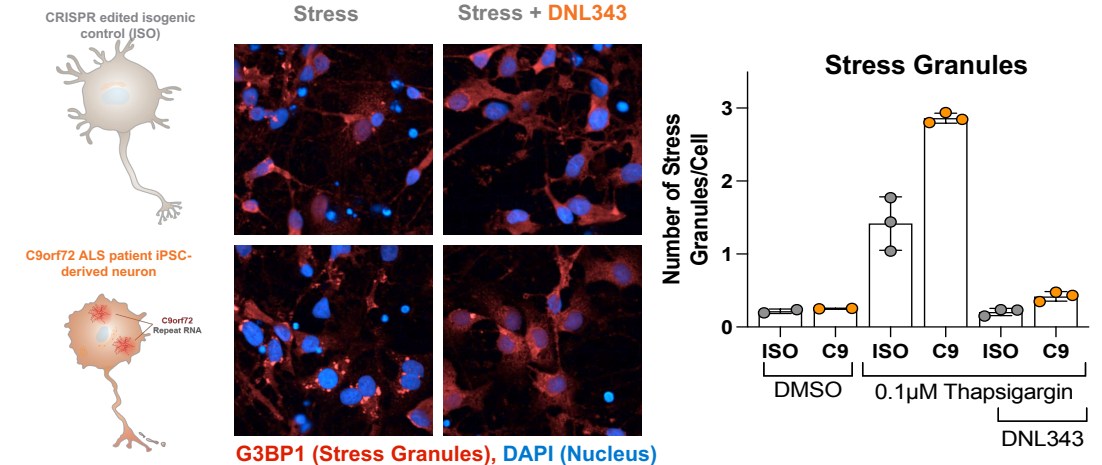
TDP-43: transactive response DNA binding protein 43 kDa; ATF4: Activating Transcription Factor 4; CHAC1: ChaC Glutathione Specific Gamma-Glutamylcyclotransferase 1

DNL343 EFFECTS IN NEURONS AND IPSC-DERIVED NEURONS

DNL343 EFFECTS IN NEURONS



DNL343 EFFECTS IN IPSC-DERIVED NEURONS



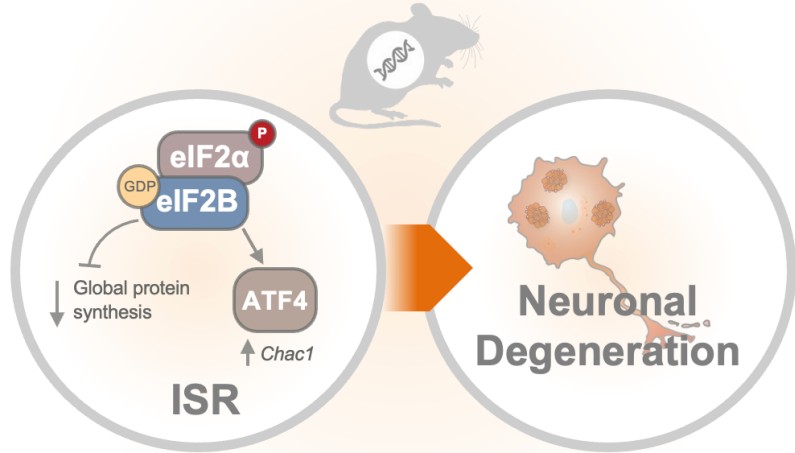
C9orf72 patient-derived neurons have increased stress granules.
DNL343 prevents this effect.

IN VITRO STUDY DESIGN

- GFP-TDP43^{ΔNLS, M337V}/mCherry-G3BP1-expressing H4 cells were treated with sodium arsenite for 1h followed by addition of 1 μM DNL343 or DMSO. Cells were imaged every 5 mins and the number of GFP⁺ puncta were quantified
- Forebrain neurons were differentiated from C9orf72-repeat containing patient iPSCs or isogenic control and matured for 2 weeks. Cells were pretreated with either DMSO or 1 μM DNL343 for 30 min followed by 2h thapsigargin treatment. Cells were then fixed and stained for G3BP1 and stress granules were quantified.

DNL343 EFFECTS IN NEURONS AND IPSC-DERIVED NEURONS

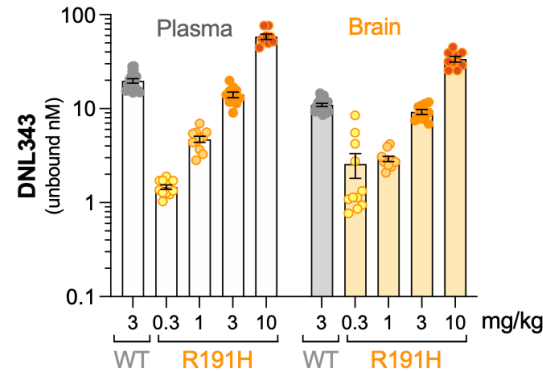
THE eIF2Bε R191H MODEL



The EIF2Bε **R191H** mouse models the consequences of ISR activation & is an ideal model to test DNL343 mechanism of action

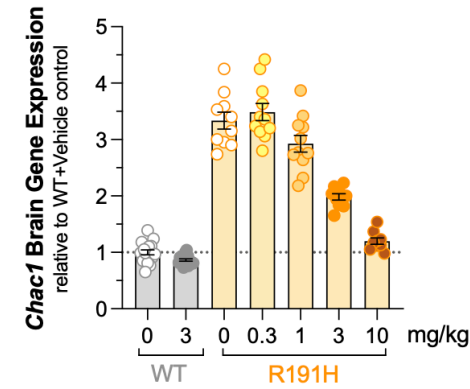
EIF2α, eIF2B, eukaryotic translation initiation factor 2α and 2B respectively; ISR, integrated stress response

DNL343 EXPOSURE



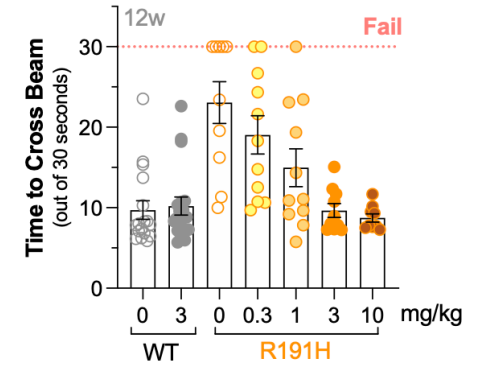
DNL343 achieved comparable exposure in the brain & plasma

ISR PATHWAY



Dose dependent modulation of brain ISR gene marker

MOTOR FUNCTION



Dose dependent restoration of motor function

DNL343 is BBB penetrant and achieved CNS pathway modulation in association with functional correction in an in vivo model

IN VIVO STUDY DESIGN

Wild-type and R191H mice self-administered chow-formulated DNL343 (*ad libitum*) for 13 weeks at doses ranging from 0.3 to 10 mg/kg daily, which led to a dose-dependent increase in exposures in the plasma and brain. Pathway modulation and functional effects were evaluated at the end of the dosing period. Data are presented as mean +/- SEM

DNL343 (eIF2B ACTIVATOR): INHIBITING THE ISR PATHWAY IN ALS

By inhibiting the ISR pathway, DNL343 is intended to prevent or slow ALS progression

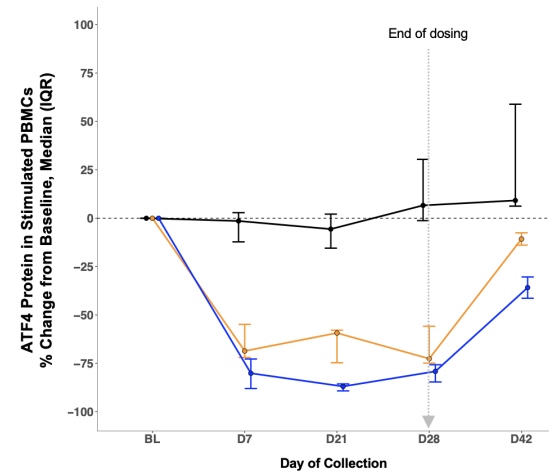
- ALS is a fatal neurodegenerative disease with TDP-43 inclusion pathology in 95% of patients
- Chronic activation of the integrated stress response (ISR) may contribute to ALS
- DNL343 is a small molecule that activates eIF2B, a key ISR regulator
- DNL343 inhibits ISR stress granule formation in cellular models
- DNL343 promotes neuroprotection in animal models

eIF2B=eukaryotic initiation factor 2B; ISR=integrated stress response; ALS=amyotrophic lateral sclerosis; TDP-43=TAR DNA-binding protein 43

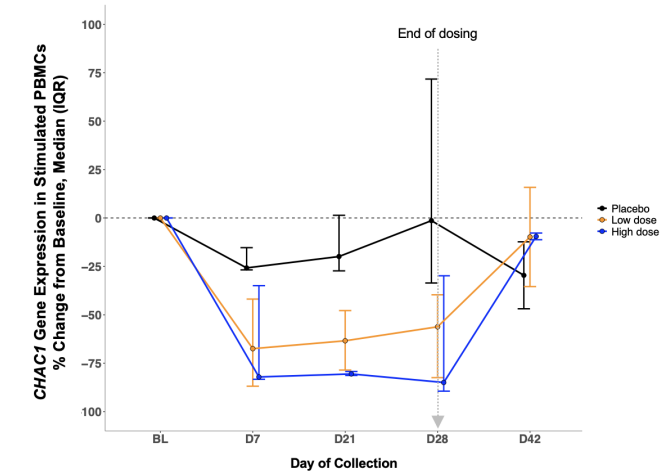
28-day dosing with DNL343 reduced ISR biomarkers in blood samples* from ALS patients (Phase 1b)



ATF4 protein levels



CHAC1 gene expression



*Fresh PBMCs were collected and stimulated ex vivo for each time point indicated for a subset of patients (per dose group: n=5-7 through day 28 and 2-3 for day 42). Experiments using cryopreserved PBMCs were also performed and showed similar results.

Dosing with DNL343 in Phase 2/3 HEALEY Platform Trial in ALS initiated May 2023

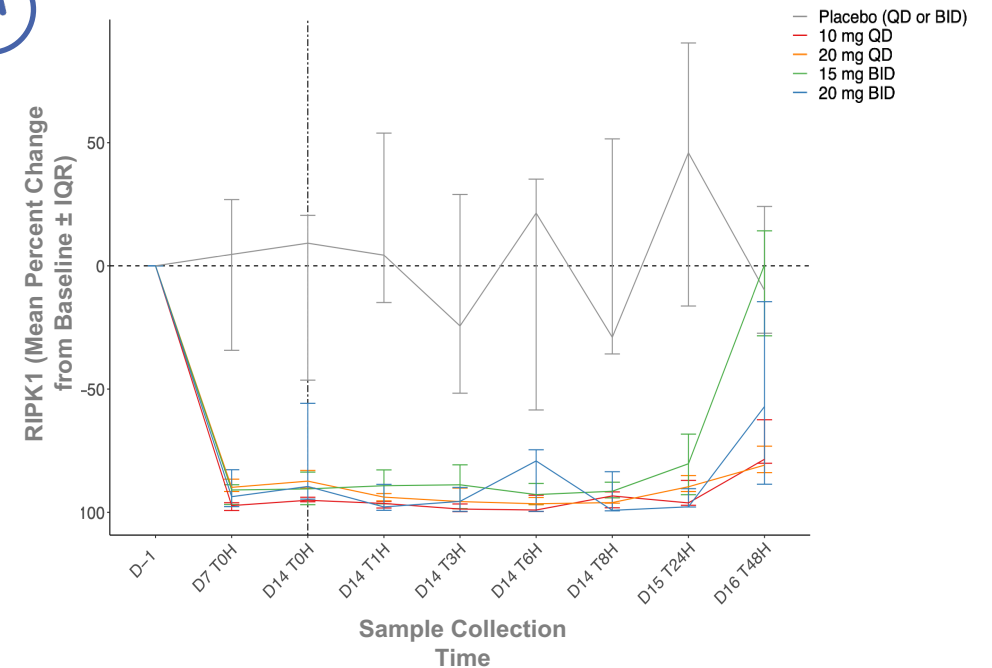
RIPK1 INHIBITORS: TARGETING INFLAMMATION AND CELL DEATH

RIPK1 is a critical signaling protein in a canonical inflammatory and cell death pathway

- Increased RIPK1 activity drives neuroinflammation and cell necroptosis and contributes to neurodegeneration
- RIPK1 inhibition achieved beneficial effects in preclinical models of ALS, multiple sclerosis and other diseases
- Denali and Sanofi have a strategic collaboration to develop and commercialize RIPK1 inhibitors
- Robust target engagement goals and safety goals achieved in Phase 1 studies for SAR443820 (CNS penetrant) and SAR443122 (peripherally restricted)

RIPK1= receptor-interacting serine/threonine-protein kinase 1; ALS=amyotrophic lateral sclerosis; MS=multiple sclerosis; UC=ulcerative colitis

93% to 99% RIPK1 inhibition achieved in Phase 1 after multiple doses of SAR443820*



*Range of maximum median inhibition of pS166-RIPK1 levels in blood cells from HVs in the Phase 1 study

**Sanofi is conducting three Phase 2 studies:
SAR443820 in ALS and MS + SAR443122 in UC**

OUR PRIORITIES

1 Clinical Execution

- 4 late-stage programs enrolling studies in MPS II, 2x ALS, and PD
- Multiple earlier-stage trials designed for biomarker PoC
- Expansion of clinical operations and medical affairs in Europe
- Building out clinical manufacturing capabilities

2 TV Expansion

- Clinical data from 3 TV-platform enabled programs
- Fourth TV-enabled program advancing towards clinical testing
- Selected OTV targets provides broad range of opportunities
- Expand TV platform potential with additional BBB transporter

3 Commercial Readiness

- Define go-to-market strategies in the US and key global markets
- Outreach to patients and communities in MPS II and ALS to understand unmet needs
- Establish critical medical affairs and commercial capabilities to prepare for early filing scenarios

TV=Transport Vehicle; OTV=Oligonucleotide Transport Vehicle; MPS= mucopolysaccharidoses; ALS=amyotrophic lateral sclerosis; PD=Parkinson's disease; PoC=proof of concept

\$1.19B in cash and investments (as of 6/30/23)

OUR PURPOSE: **DEFEAT DEGENERATION**

Thank you to all those who are part of Denali's purpose,
especially our patients and their families



**LYSOSOMAL STORAGE
DISEASE**



**RARE NEURODEGENERATIVE
DISEASES (ALS, FTD)**



**PARKINSON'S
DISEASE**



**ALZHEIMER'S
DISEASE**



Denali

The name captures the formidable challenges in fighting neurodegenerative diseases but also the unprecedented opportunities enabled by new scientific insights and technologies. With a relentlessly committed team and rigorous effort, breakthroughs appear to be within reach.

A photograph of the Golden Gate Bridge in San Francisco, viewed from a high vantage point on a rocky, vegetated cliff. The bridge's iconic red-orange towers and suspension cables are prominent, extending across the frame. The bridge deck is visible, with some vehicles. The water below is a deep blue-green, and a thick layer of white fog or mist hangs over the water and the lower part of the bridge. The sky is overcast and grey. The text "THANK YOU" is superimposed in the center of the image in a white, sans-serif font.

THANK YOU