

Alector Company Overview

September 2022

Forward-Looking Statement

This presentation contains forward-looking statements that involve substantial risks and uncertainties. All statements other than statements of historical facts contained in this presentation are forward-looking statements. In some cases, you can identify forward-looking statements by terminology such as "anticipate," "believe," "continue," "could," "estimate," "expect," "intend," "may," "plan," "potentially," "predict," "should," "will" or the negative of these terms or other similar expressions. Forward-looking statements contained in this presentation also include, but are not limited to, statements regarding: our future financial condition, results of operations, business strategy and plans, plans, timelines and expectations related to our product candidates and our other clinical programs, including with respect to the availability of data, the initiation of future clinical trials and plans and expectations regarding planned regulatory filings with respect to such programs; and objectives of management for future operations, as well as statements regarding industry trends.

We have based these forward-looking statements largely on our current expectations and projections about future events and trends that we believe may affect our financial condition, results of operations, business strategy and financial needs. These forward-looking statements are subject to a number of risks, uncertainties and assumptions, including, among other things: Alector's plans relating to the development and manufacturing of its product candidates and research programs; the ability of Alector, Inc.'s ("Alector") clinical trials to demonstrate safety and efficacy of its product candidates, and other positive results; the timing and focus of Alector's trategic collaborations with third parties and Alector's ability to expende potential benefits of strategic collaborations with third parties and Alector's setimates of the number of patients in the United States who suffer from the diseases it is targeting and the number of patients that will enroll in its clinical trials; the size of the market opportunity for Alector's product candidates in each of the diseases it is targeting; Alector's ability to expand its product candidates, such as orphan drug designation, for its product candidates for various diseases; Alector's ability to obtain and maintain regulatory approval of regulatory filings and approvals, including Adettor's continued reliance on third parties to conduct additional clinical trials of its product candidates, and for the manufacture of its product candidates for preclinical studies for preclinical studies for preclinical studies and clinical trials; and ther pursidictions; Alector's continued reliance on third parties to conduct additional clinical trials of its product candidates, and for the manufacture of its product candidates for preclinical studies for preclinical studies for preclinical studies and clinical trials; and the other jurisdictions; Alector's continued reliance on third parties to conduct additional clinical trials of its product candidates, and for the manufacture of its

This presentation also contains results based on data from our clinical trials. These clinical trials are ongoing and this presentation does not speak to, and you should make no assumptions about, any additional data. In addition, the information we have chosen to publicly disclose regarding our product candidates has been selected from a more extensive amount of available information. You or others may not agree with what we determine is the material or otherwise appropriate information to include in our disclosure, and any information we determine not to disclose may ultimately be deemed significant with respect to future decisions, conclusions, views, activities or otherwise. If the initial data that we report differ from updated, late, final or actual results, or if others, including regulatory authorities, disagree with the conclusions reached, our ability to obtain approval for, and commercialize our other product candidates may be harmed, which could harm our business, financial condition, results of operations and prospects.

Except as required by law, we undertake no obligation to update any statements in this presentation for any reason after the date of this presentation.

We have filed Current Reports on Form 8-K, Quarterly Reports on Form 10-Q, Annual Reports on Form 10-K, and other documents with the SEC. You should read these documents for more complete information about us. You may obtain these documents for free by visiting EDGAR on the SEC website at www.sec.gov.

This presentation discusses certain investigational therapeutic agents, which have not yet been approved for marketing by the U.S. Food and Drug Administration. No representation is made as to the safety or effectiveness of our product candidate for the therapeutic use for which it is being studied.

This presentation contains statistical data based on independent industry publications or other publicly available information, as well as other information based on our internal sources. We have not independently verified the accuracy or completeness of the data contained in these industry publications and other publicly available information. Accordingly, we make no representations as to the accuracy or completeness of that data.



Translating Scientific Insights into a Broad Portfolio of First-in-Class Programs

NOVEL APPROACH	Founded to pioneer a new field of research: Immuno-neurology	Informed by neuroscience , human genetics and immunology	Substantial IP portfolio established: <i>20 issued patents,</i> <i>450+ patent applications</i>
MULTIPLE CLINICAL TRIALS	Phase III Clinical Program for FTD-PGRN	Clinical Programs for AD, FTD-GRN, FTD- C9ORF72, ALS	Pre-Clinical Programs for AD, PD, Solid tumors
WORLD CLASS PARTNERS	\$700M upfront \$1.5B+ milestone 50-50 U.S. profit share Tiered double-digit royalties	\$20M ed \$986M r	upfront payment quity investment nilestone payments 0-50 profit share
STRONG FINANCIALS	\$809 MILLION IN CASH		
alector ⁻	FTD = Frontotemporal dementia, PD = Parkinson's Dise	ase, AD = Alzheimer's Disease, ALS = Amyotrophic lateral	sclerosis Proprietary and Confidential Property of Alector 3

Experienced Leadership and Advisors Guide Clinical and Corporate Execution

MANAGEMENT



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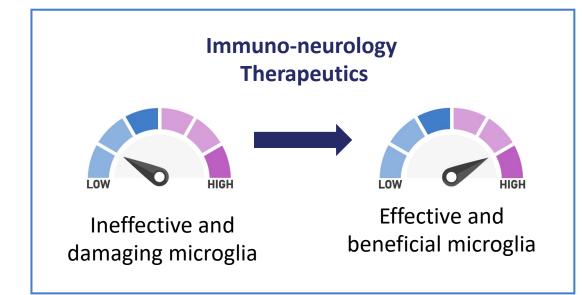
SCIENTIFIC ADVISORY BOARD

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John Maraganore appointed as Strategic Advisor

Immuno-Neurology: Alector's Therapeutic Strategy for Degenerative Brain Disorders

Recruiting microglia, the brain's immune system, to potentially cure neurodegeneration



Using the immuno-oncology concept of harnessing the immune system as a broad, effective and long-lasting therapy

Multiple programs are in or entering the clinic for multiple neurodegenerative diseases

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Human Genetics

Develop drugs targeting risk genes for neurodegeneration to functionally enhance protective functions of these risk genes

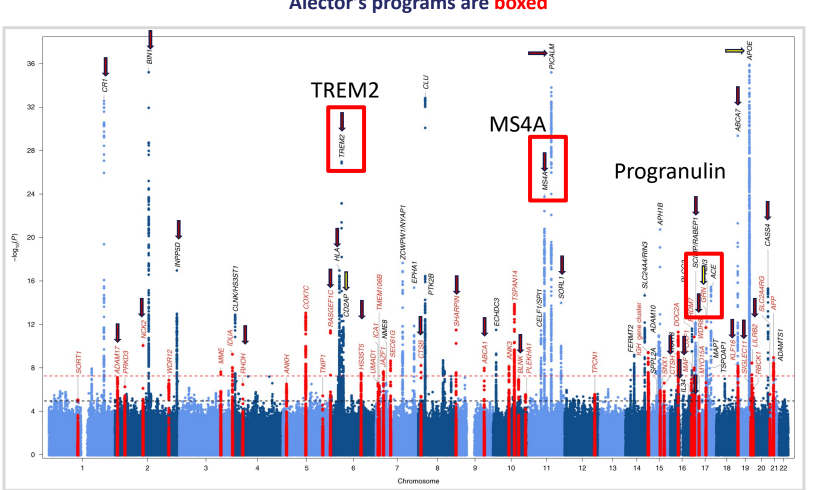
Immunology Target checkpoint regulators on microglia and harness microglia as broad therapy for neurodegeneration

Neuroscience

Rejuvenate microglia and harness their physiological role as guardians of brain health

Immuno-neurology: targeting the brain immune system as a novel therapeutic strategy for dementia and neurodegeneration

Genetic Rationale for Immuno-Neurology

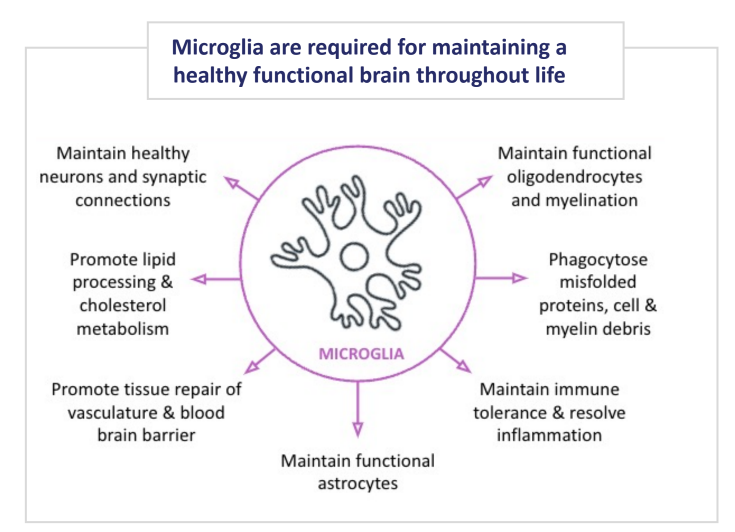


Many AD risk genes are regulators of the microglia brain immune cells (Arrows in black) Alector's programs are boxed

Bellenguez C et al. Nat Genet. 2022; 54(4): 412–36.



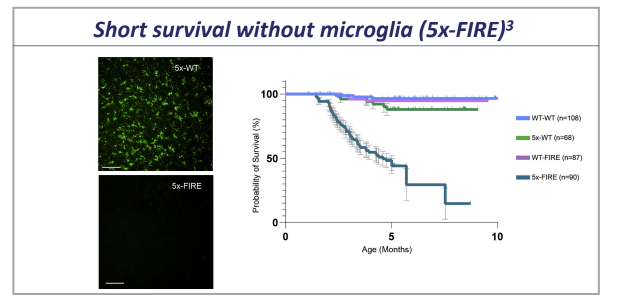
Biological Rationale for Immuno-Neurology: The Microglia Brain Immune System is Essential for Brain Function and Health



Targeting microglia immune checkpoints and harnessing microglia to cure neurodegeneration

Microglia Are Essential for Brain Health in Mouse Models

- "Microglia jointly degrade fibrillar alpha-synuclein cargo by distribution through tunneling nanotubes"¹
- "Negative feedback control of neuronal activity by microglia"²
- Absence of microglia in AD mice lead to cerebral amyloid angiopathy, hemorrhages, calcification, and lethality³
- Transplantation of microglia reverses these pathological changes ³



or ¹Scheiblich H et al. *Cell* 2021;184(20):5089-106.e21. ²Badimon A et al. *Nature* 2020;586(7829):417-23. ³Shabestari SK et al. *Cell Rep.* 2022;39(11):110961.

Immuno-neurology: targeting the brain immune system as a novel therapeutic strategy for dementia and neurodegeneration

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Microglia Are Essential for Brain Health in Humans

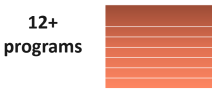
Loss of microglia due to CSF1R mutations neurodegeneration "Adult- Onset Leukoencephalopathy"		Patients experience range of psychiatric, neurocognitive, and motor symptoms; Average age of onset is ~43		
Rapid brain tissue loss	~ 6 Year Survival Ra	te Rapid Cognitive Decline		
tient VI 2442+1G>T) Years before onset yo Year after onset	(b) 1.0 0.8 0.6 0.4 0.4 0.2 0.0 0.0 0.0	 and an analysis be a patient A construction and a patient A be a patient B be a patient C construction constructin construction construction		

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Portfolio of Product Candidates Targeting Genetic Causes of Neurodegeneration as well as Promising Innate Immune System Targets for Oncology

	TARGET	CANDIDATE	RESEARCH	PRECLINICAL	PHASE 1	PHASE 2	PHASE 3	COMMERCIAL RIGHTS
		AL001	FTD-GRN				>	gsk 🗤
Progranulin	DCDN	AL001	FTD- <i>C9orf</i> 72			>		gsk 📊
Franchise	PGRN		ALS*			>		gsk 🕕
		AL101	Healthy volunteers for AD and PD >		>			gsk 🕕
	TREM2	AL002	Alzheimer's disease			>		abbvie 🚛
Alzheimer Programs		AL044	Alzheimer's disease	>				
AL044			Orphan neuro indicati	on >				
Oncology	SIRP-alpha	AL008	Solid tumors	>				
Programs	Multi-Siglec	AL009	Solid tumors >					

Target indications include AD, PD, FTD, MS & cancer



IP portfolio contains 50+ patent application families, which include 20 issued patents and >450 pending patent applications directed to more than 20 targets and/or technologies



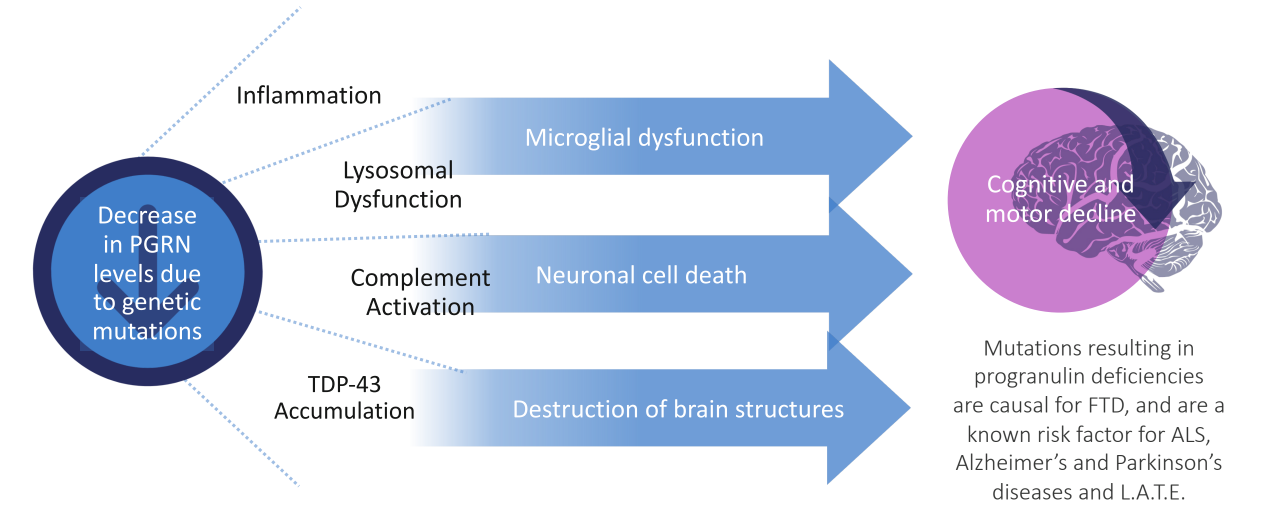
AD = Alzheimer's disease PD = Parkinson's disease FTD = Frontotemporal dementia ALS = Amyotrophic lateral sclerosis MS = Multiple sclerosis

*In partnership with GSK, the company made a strategic, non-safety related decision to close enrollment in the ALS-C9orf72 Phase 2a biomarker trial and is currently evaluating plans for a potential Phase 2b study for patients with all forms of ALS, including the C9orf72 mutation.

Progranulin Franchise Programs AL001 / AL101

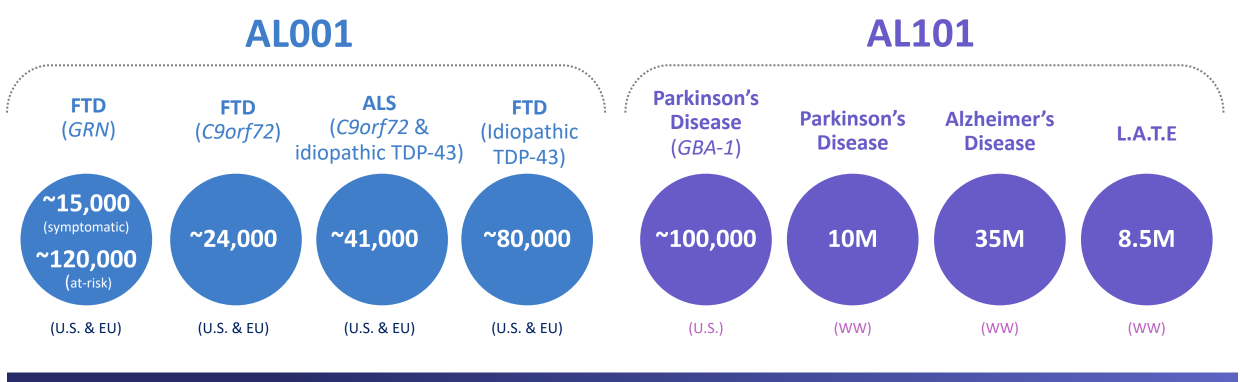


The Role of Progranulin in Neurodegeneration





Broad Therapeutic Potential Grounded in Genetic Evidence and Animal Models



GENETIC EVIDENCE

Known Risk Factor/ Positive Correlation

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Causal

FTD = Frontotemporal dementia ALS = Amyotrophic lateral sclerosis L.A.T.E. = Limbic-predominant age-associated TDP43 encephalopathy

AL001 and AL101: Targeting Progranulin to Restore Function of Microglia

MECHANISM

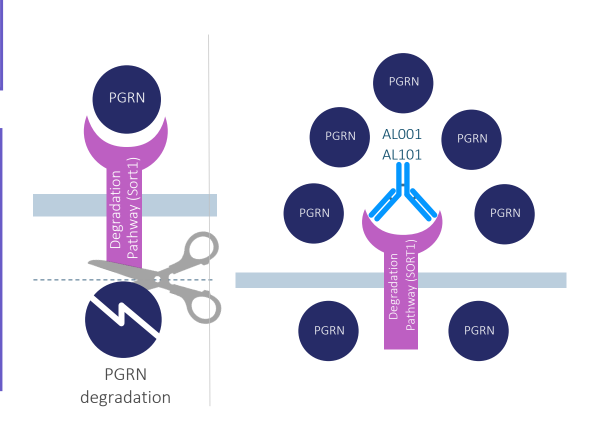
Increases the half-life of PGRN by blocking Sortilin, a degradation receptor, in order to restore PGRN to normal levels

AL001 STATUS

- Orphan Drug and Fast Track Designation
- Phase 1 studies in healthy volunteers complete
- Data presented from ongoing Phase 2 study in FTD-GRN and FTD-C9orf72
- Currently evaluating study design options for Phase 2b study in patients with all forms of ALS, including the *C9orf72* mutation
- Pivotal Phase 3 study in FTD-GRN actively enrolling

AL101 STATUS

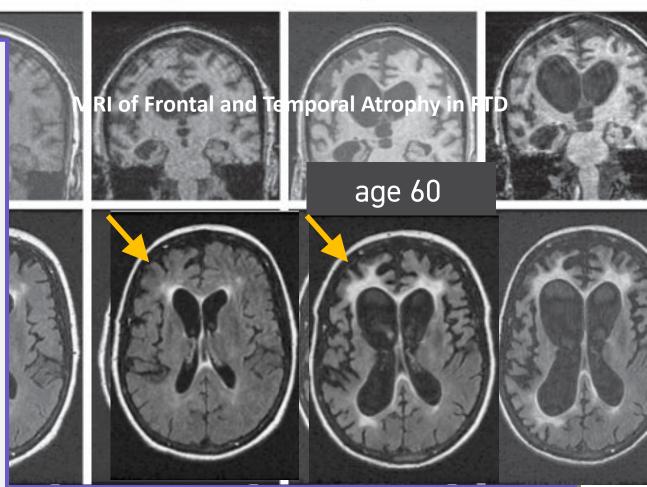
• Ongoing Phase 1 study in healthy volunteers





Frontotemporal Dementia: A Rapidly Progressive Form of Dementia: With No
Evaluation 3Current TreatmentAge 57Age 58Age 59Age 60Age 61

- Most common form of dementia under the age of 60
- Patients present with compulsive behavior, lack of restraint, apathy, anxiety, and aphasia
- Life expectancy after diagnosis is 7 10 years
- 15,000 symptomatic + ~120,000 at-risk people with progranulin mutations (FTD-GRN) in the U.S. and E.U.
 - FTD-GRN caused by coding mutations in progranulin
 - Lead to a complete loss of function in the mutated gene





INFRONT-2: Phase 2 in Frontotemporal Dementia Populations



Twelve-month biomarker and clinical data presented at 2021 AAIC and CTAD conferences from the symptomatic FTD-*GRN* cohort



- Asymptomatic and symptomatic FTD-GRN enrollment closed; FTD-C9orf72 cohort currently enrolling
- 2. CDR® plus NACC FTLD-SB: Clinical Dementia Rating (CDR) dementia staging instrument plus National Alzheimer's Coordinating Center (NACC) behavior and language domains frontotemporal lobar degeneration (FTLD) sum of boxes (SB)

Latozinemab Impacts Key Markers of the Disease Cascade in Symptomatic FTD-GRN Patients

Key biomarkers and clinical outcome assessments reflect underlying disease activity in FTD-GRN patients

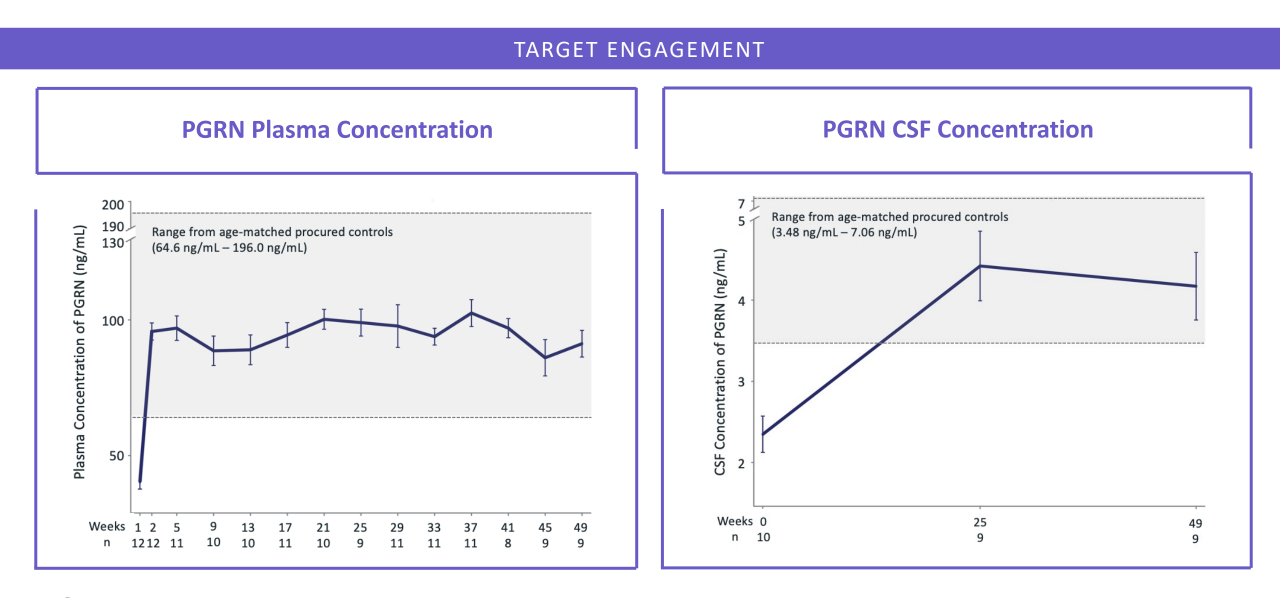
TARGET ENGAGEMENT	BIOMARKERS OF DISEASE ACTIVITY				CLINICAL BENEFIT
PGRN (plasma and CSF)	Lysosomal dysfunction	Complement activation	Astrogliosis	Neuronal health	Clinical Outcome Assessments
PGRN	e.g. CTSD, LAMP1	e.g. C1QB	GFAP	NfL	CDR [®] plus NACC FTLD-SB
> 50% reduction in PGRN levels causal for FTD	Dysfunctional lysosomes are hallmark of FTD- <i>GRN</i>	Pathological increases in complement proteins in FTD correlate with cognitive decline	GFAP is increased in conditions characterized by astrogliosis	NfL is a measure of axonal damage	FDA approvable endpoint for measuring clinical decline in FTD

CTSD = cathepsin D; LAMP1 = lysosomal associated membrane protein 1; C1QB = complement C1q B chain; GFAP = glial fibrillary acidic protein NfL = neurofilament light chain;



CDR[®] plus NACC FTLD-SB: Clinical Dementia Rating (CDR) dementia staging instrument plus National Alzheimer's Coordinating Center (NACC) behavior and language domains frontotemporal lobar degeneration (FTLD) sum of boxes (SB)

INFRONT-2: AL001 Restores PGRN in Plasma and CSF to Normal Levels



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 Data cut-off June 15, 2021

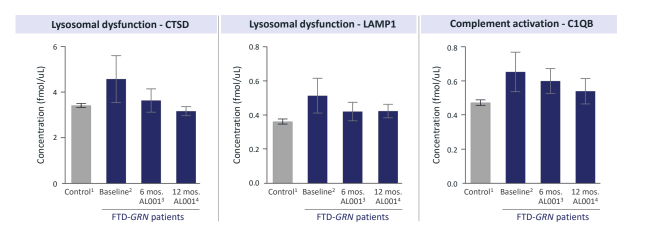
 Mean +/- SEM

INFRONT-2: Latozinemab Demonstrated Consistent Effects on Disease Biomarkers

FLUID BIOMARKERS OF DISEASE ACTIVITY

Symptomatic FTD-GRN patients at 12 months

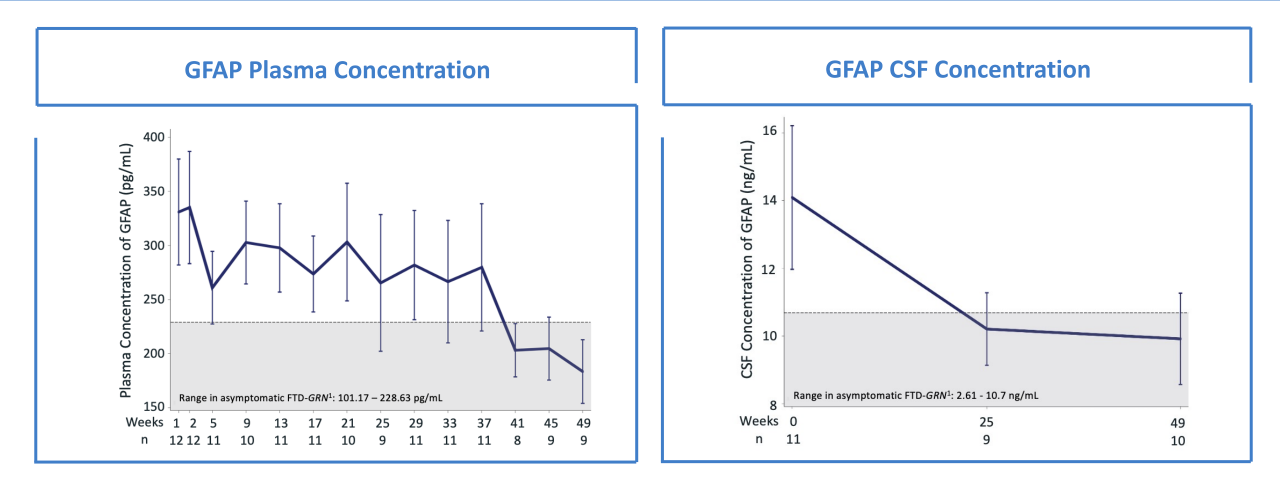
Normalization of lysosomal and inflammatory biomarkers



Markers	Latozinemab Baseline (N=9)	Latozinemab 6 months (N=8)	Latozinemab 12 months (N=8)	Age- matched procured control (N=44)
CTSD (fm/µL)	5.2 (1.16)	3.8 (0.57)	3.1 (0.21)	3.4 (0.08)
LAMP1 (fm/µL)	0.6 (0.12)	0.4 (0.06)	0.4 (0.043)	0.4 (0.01)
C1QB (fm/µL)	0.7 (0.12)	0.6 (0.07)	0.5 (0.02)	0.5 (0.02)

INFRONT-2: AL001 Treatment Decreases Glial Fibrillary Acidic Protein (GFAP) Levels Towards Range Seen in Asymptomatic FTD-GRN

BIOMARKERS OF DISEASE ACTIVITY – ASTROGLIOSIS



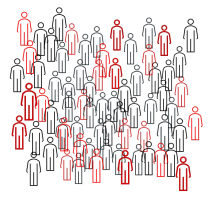


1. Range is of baseline GFAP levels in asymptomatic FTD-GRN patients enrolled in INFRONT-2

INFRONT-2: A two-step Matching Strategy to Eliminate Potential Confounding Factors in Constructing the GENFI2 Historical Control Cohort



GENFI2 FTD-GRN participants with at least one post-baseline CDR[®] plus NACC FTLD-SB (n=102)

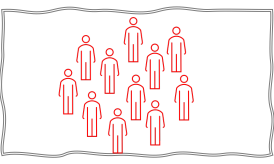


Propensity score matching by baseline CDR[®] plus NACC FTLD-SB Step 2 Potential GENFI2 matches using propensity score matching based on CDR[®] plus NACC FTLD-SB (n=25) Blin



Blinded clinical adjudication matching by NfL at baseline, age, diagnosis and gender

<u>GENFI2 matched historical</u> <u>control cohort (n=10)</u>



- Propensity scores were computed using a logistic regression including the most important covariate, cognition, measured at baseline using the CDR[®] plus NACC FTLD-SB
- Matching was done by comparing the logit propensity score

- To further increase the chances that the matched historical control cohort would mimic a placebo group in a randomized experiment, clinical adjudication of secondary covariates, including NfL at baseline, age, diagnosis and gender were used to refine and construct the final matched historical control cohort
- This step was done on a blinded basis without knowing the progression rate.



INFRONT-2: Contextualizing vMRI and Clinical Results with GENFI2 Matched Controls

INFRONT-2 vMRI and clinical results compared against comparable, matched GENFI2 controls

- Comparable baseline characteristics established between INFRONT-2 FTD-GRN symptomatic cohort and GENFI2 controls in two-steps:
- Propensity score matching¹ based on CDR[®] plus NACC FTLD-SB at baseline
- Matching refined by age, gender, baseline plasma NfL levels, and FTD disease variant at baseline²

Baseline characteristics		INFRONT-2 (N=12)	GENFI2 Matched Controls (N=10)	
CDR [®] plus NACC FTLD-SB Mean (SD)		5.9 (3.74)	5.2 (3.60)	
	Min, Max	0.5, 11	0.5, 11.5	
AGE (Years)	Mean (SD)	63.2 (9.71)	62.1 (6.74)	
	Min, Max	49, 79	52, 72	
GENDER	Male	8 (67%)	3 (30%)	
PLASMA NfL (pg/mL)	Ν	12	9	
	Mean (SD)	62.8 (47.00)	40.3 (27.28)	
	Min, Max	11.2, 148.8	9.3, 99.9	
FTD DISEASE VARIANT	bvFTD	5 (42%)	5 (50%)	
	PPA	3 (25%)	3 (30%)	
	Both	3 (25%)	0	
	Other	1 (8%)	1 (10%)	

GENFI = The Genetic Frontotemporal Initiative

GENFI2 refers to the longitudinal FTD registry dataset

1. Propensity score matching is a well-established statistical method intended to mimic randomization

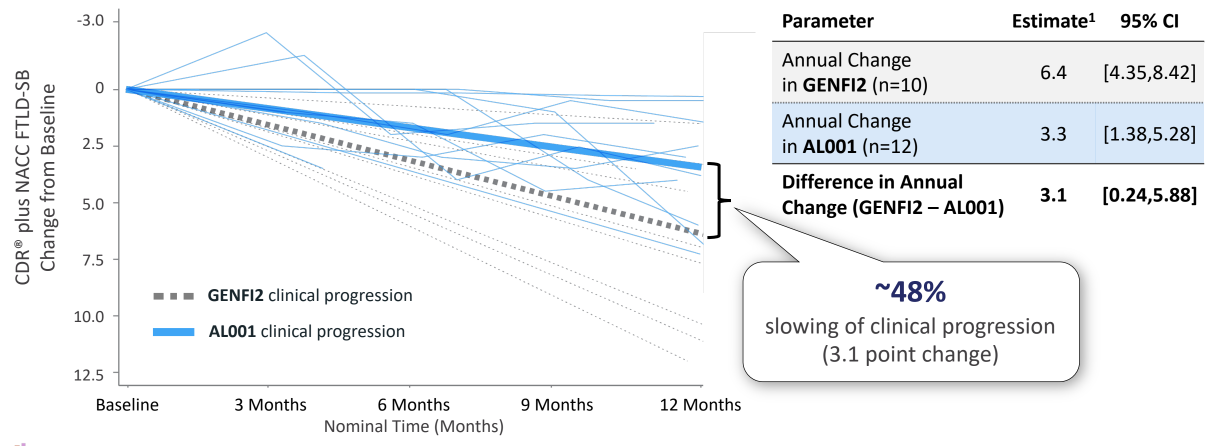


Latozinemab-Treated FTD-GRN Parcipipants Experience a ~48% Annual Delay in Disease Progression Compared to Matched Historical Controls

Similar findings of ~54% delay in disease progression for C9-orf72 FTD patients: Latozinemab was well tolerated in INFRONT-2

CLINICAL BENEFIT

CDR® plus NACC FTLD-SB



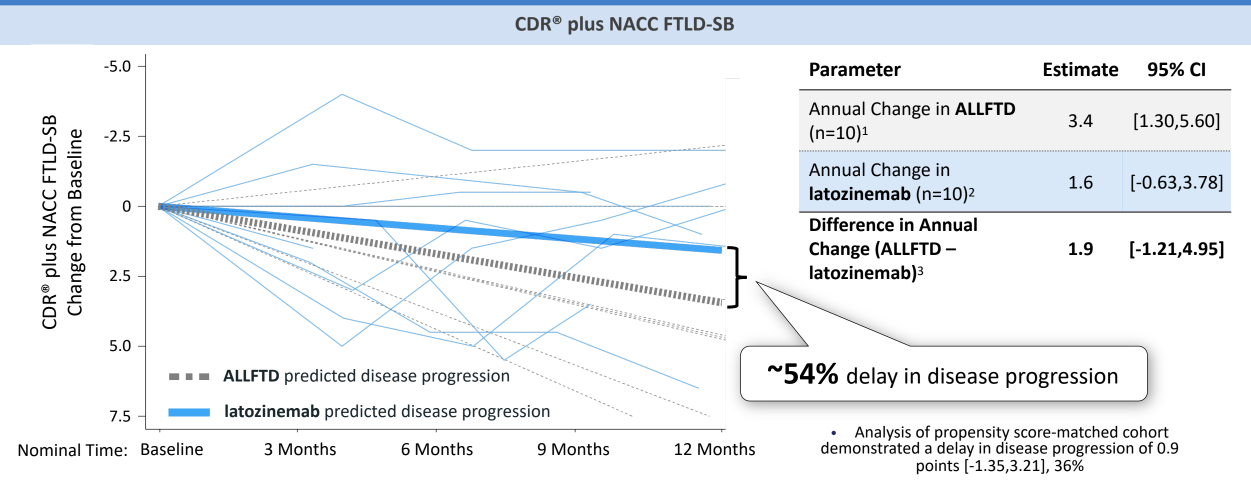
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Random Coefficient Model with Repeated Measurements including baseline & all available post-baseline measurements up to 12 months. Data cut-off Sep 8, 20: Phase 2 data presented at CTAD 2021 and ADPD 2022 NCT03987295 Immuno-neurology: targeting the brain immune system as a novel therapeutic strategy for dementia and neurodegeneration

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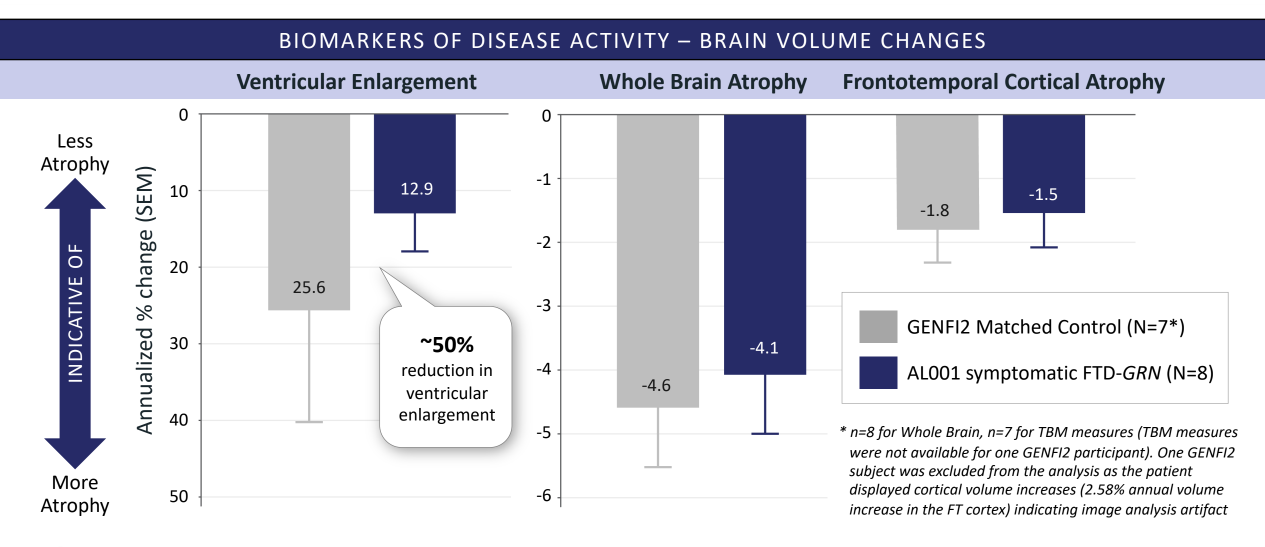
When Compared to the ALLFTD Matched Historical Controls, Latozinemab-Treated FTD-C9orf72 Participants Experience a ~54% Annual Delay in Disease Progression

CLINICAL BENEFIT



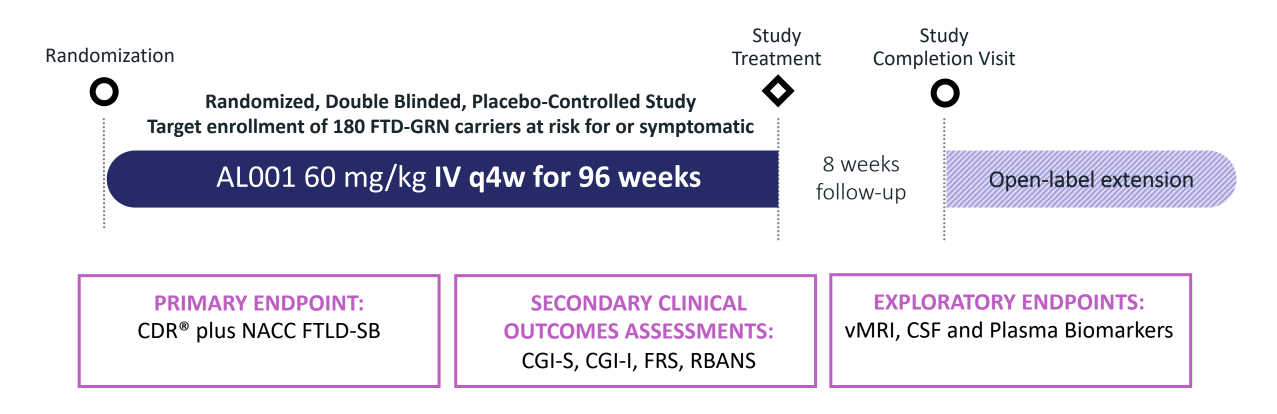


INFRONT-2: vMRI Data Suggest Slowing of Ventricular Enlargement and Brain Atrophy in AL001-Treated Patients vs. Historic Matched Control



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Enrollment Ongoing for Pivotal INFRONT-3 Phase 3 Study of AL001



Study taking place at approximately 45 clinical centers in US, Canada, Europe and Australia

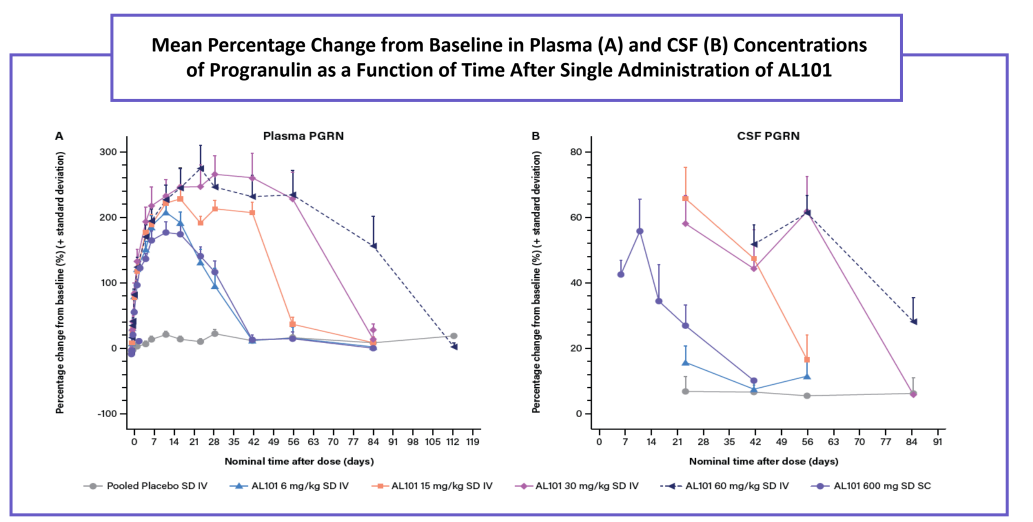
Initial data read out after 96-week treatment period



"At risk" = GRN carriers who are pre-symptomatic and meet a pre-specified NfL threshold for enrollment in the Phase 3 CDR® plus NACC FT1. LD-SB = Clinical Dementia Rating Dementia Staging Instrument PLUS National Alzheimer's Disease Coordinating Center Frontotemporal Lobar Degeneration Behavior and Language Domains Sum of Boxes; CGI-S = Clinician's Global Impression-Severity; CGI-I = Clinician's Global Impression-Improvement; FRS = Frontotemporal Dementia Rating Scale; RBANS = Repeatable Battery for the Assessment of Neuropsychological Status

AL101 Elevated Progranulin Levels in Periphery and CSF in Phase 1

Generally well tolerated in healthy volunteers; initial Phase 1 data suggest less frequent dosing may be possible



Phase 1 ongoing with additional cohorts being enrolled to evaluate subcutaneous dosing



Clinical-Stage Alzheimer's Disease Candidate: AL002



AL002: Designed to Activate TREM2 in Order to Enhance Microglia Function

RATIONALE

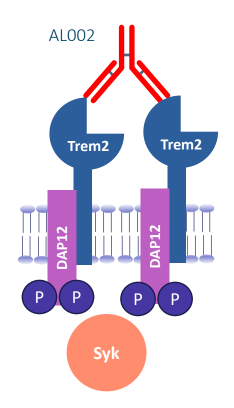
- TREM2 signaling controls critical microglial activity
- TREM2 is a prominent risk gene for Alzheimer's disease
 - Homozygous mutations cause dementia (NHD, FTD)
 - Heterozygous mutations increase risk for Alzheimer's disease by 3x
 - Ligands include APOE, an Alzheimer's risk gene

MECHANISM

 Activates TREM2 signaling to improve functionality of microglia to clear pathology and protect neurons

STATUS

- Phase 1 study complete
- INVOKE-2 Phase 2 clinical trial on-going

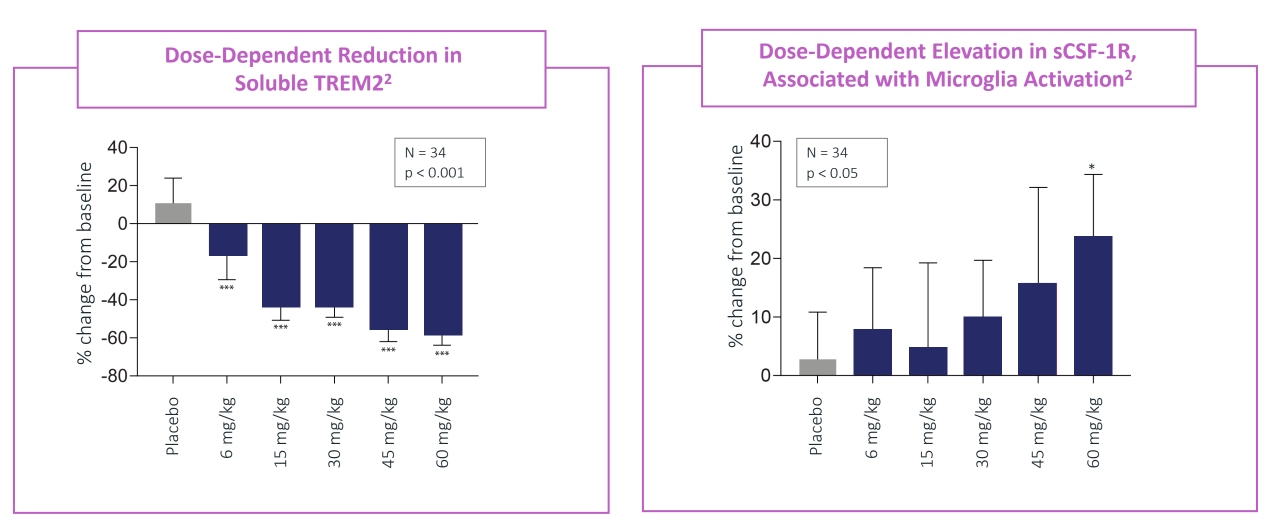


Intended to improve survival, proliferation, function of microglia



AL002 Target and Microglia Engagement Achieved in Phase 1

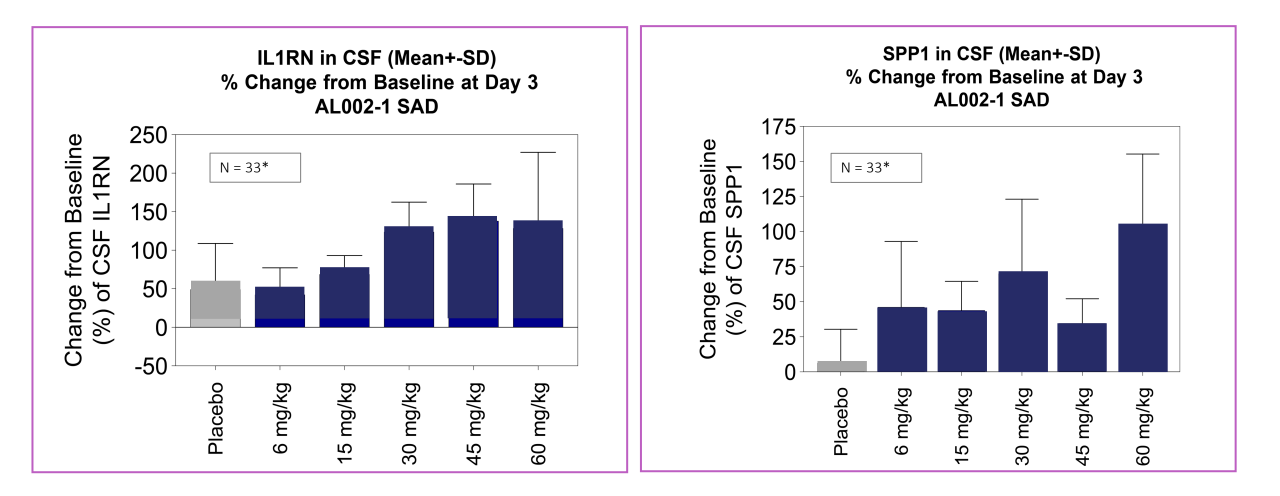
Generally well-tolerated in healthy volunteers¹



Data are presented as mean ±SD; cohort n = 6 (placebo, 6 mg/kg, 15 mg/kg, 30 mg/kg) and 5 (45 mg/kg, 60 mg/kg). ***P = 0.0001 for 6 mg/kg and P < 0.0001 for all other doses vs. pooled placebo control. *P = 0.026 at 60 mg/kg vs. pooled placebo.

¹Phase 1 data presented at AAIC 2021; NCT03635047. ²Wang S et al. *J Exp Med*. 2020;217(9):e 20200785.

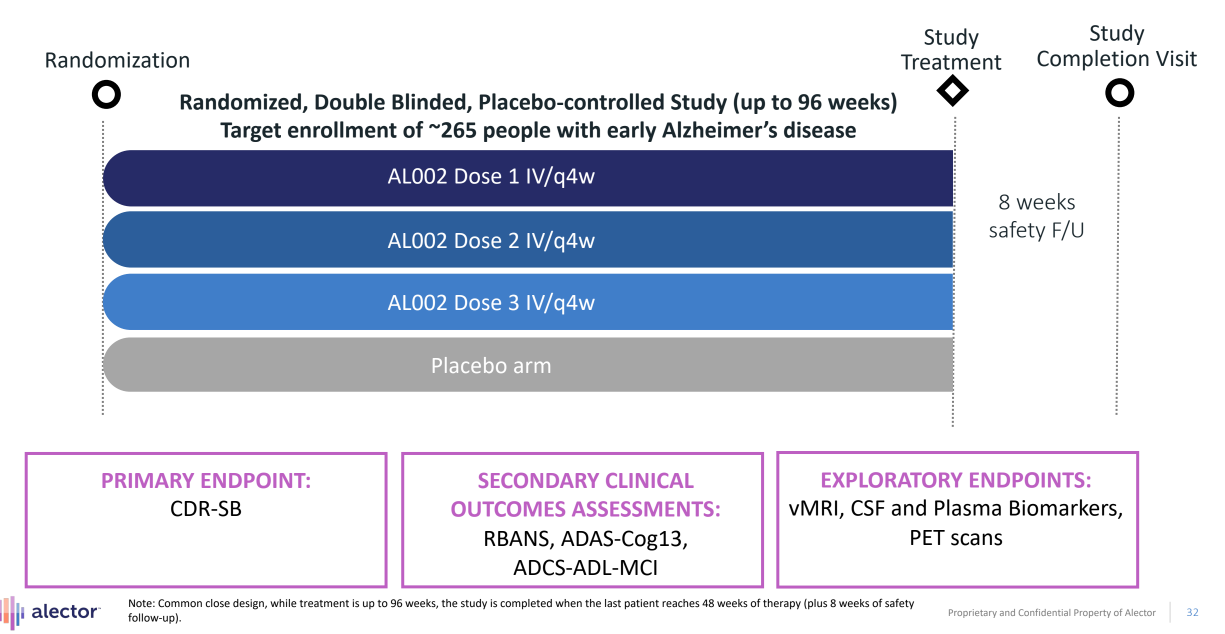
AL002 Treatment in Phase 1 Also Caused An Increase in CSF Levels of IL1RN and SPP1, Indicating Further Evidence of Microglial Activation





At doses 6mg/kg – 45 mg/kg, N=6/cohort. N=14 in the 60 mg/kg cohort. Pooled placebo N=11. *Outlier value (1280.3% above baseline at Day 3 from 1 participant in the 30 mg/kg dose group were omitted from the graph. Phase 1 data presented AAIC 2021

INVOKE-2 Phase 2 AL002 Study in Individuals with Early Alzheimer's Disease



Preclinical Program for Alzheimer's Disease: AL044



Background on AL044 Targeting a Candidate Master Regulator of Microglia

Key Features of MS4A and AL044

MS4A Member of ~22 membrane-tetra spanning 4A family, structurally related to CD20

Expressed on microglia, CNS perivascular macrophages

Modulate multiple aspects of AD disease risk, age of onset, progression and survival

AL044 our drug candidate functionally phenocopies and exceeds activities of the protective MS4A variant

AL044 Regulates the levels of key signaling systems in microglia; Trem2/sTrem2, CSF1R, Dectin1

AL044 regulate microglia , proliferation, survival migration lysosomal function, immune response and energetics, genes and/or proteins

IND filed following pre-IND alignment

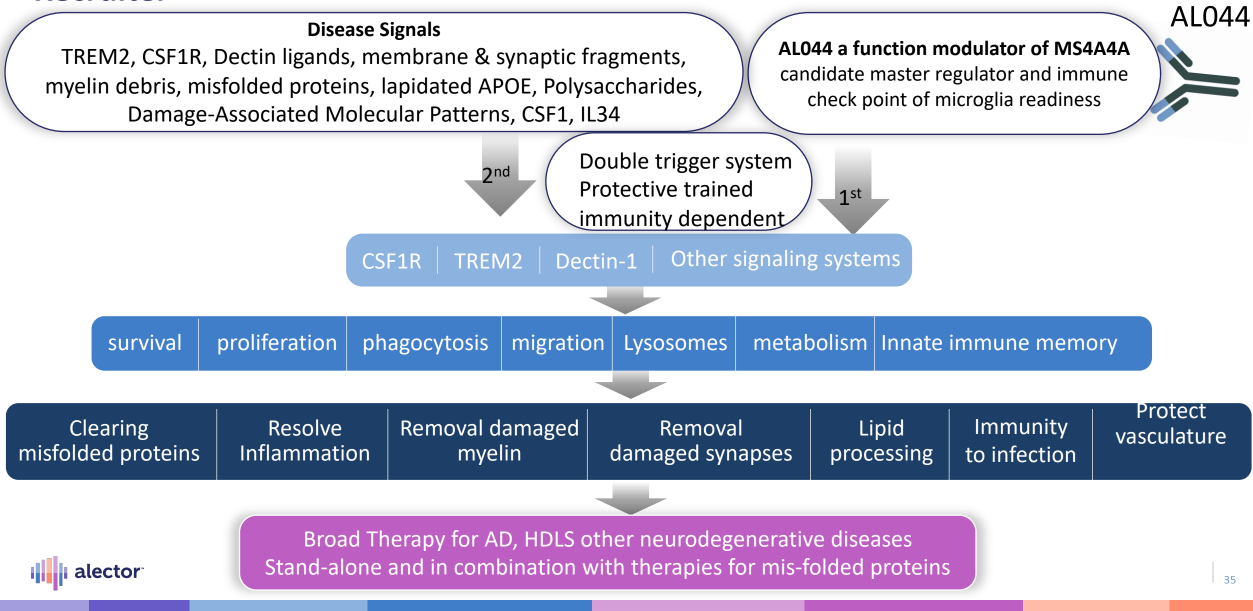
Source: doi:10.1126/scitranslmed.aau2291, Sci Transl Med, 2019.; NATURE IMMUNOLOGY | VOL 20 | AUGUST 2019 | 1012–1022; DOI: 10.1016/j.jalz.2016.06.005; Michael Ewers et al., Sci Transl Med 2019;11:eaav6221.

Effects of MS4A on AD

Protective Allele	Effects on AD	Risk Allele
	AD Risk	
	Age of onset and survival	
	Rate of cognitive decline	
	CSF Soluble TREM2	
	Aβ Plaques & CSF Tau	
	Rate of Cortical and hippocampal Shrinkage	
	Rate of Conversion from MCI to AD	
	Protective Interactions with APOE4	

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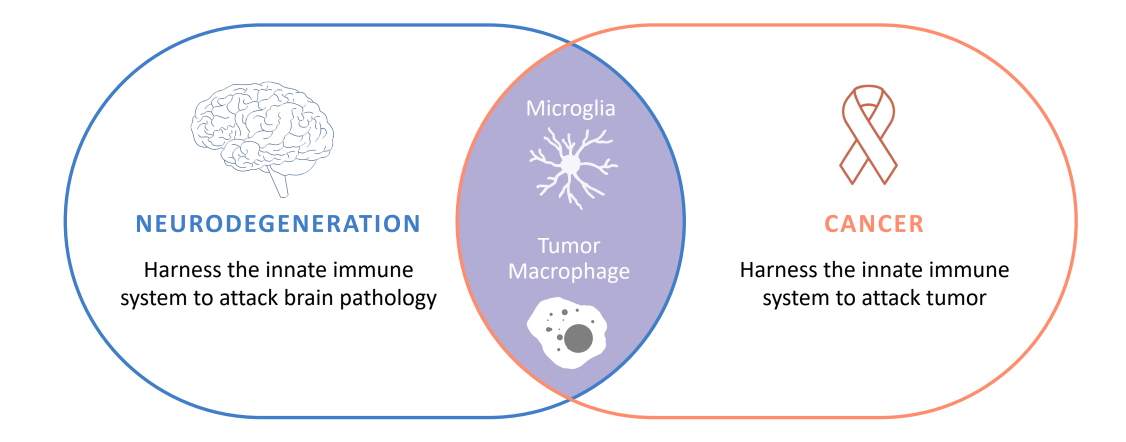
AL044 Mechanism of Action: a Context and Memory Dependent Microglia Recruiter



Alector Oncology Overview



Neurodegeneration and Cancer Converge at the Innate Immune System





AL009: Marshalling the Innate Immune System to Combat Tumor Growth

TARGET

Siglec-Sialic acid innate checkpoint pathway

SCIENTIFIC RATIONALE

Human genetics and tumor model data show Siglecs drive immune suppression

STATUS

IND Submission expected later this year

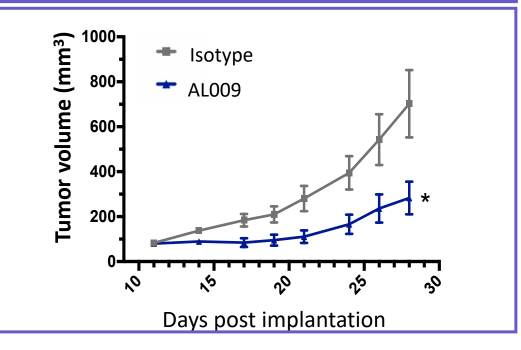
PRODUCT CANDIDATE

- Engineered Fc fusion protein
- Blocks multiple Siglecs to activate innate and adaptive immunity
- Engages activating Fcγ receptors

PRECLINICAL ACTIVITY

- Repolarizes suppressive macrophages, activates T cells and enhances ADCC/ADCP function
- Demonstrates robust anti-tumor activity in multiple models (single agent and combined with checkpoint inhibitors)

Monotherapy Activity in Breast Cancer Model





AL008: Potential Best-in-Class Dual Function SIRPα-CD47 Pathway Activator

TARGET

SIRP α - CD47 pathway

SCIENTIFIC RATIONALE

Tumors leverage pathway to hide from immune system

STATUS

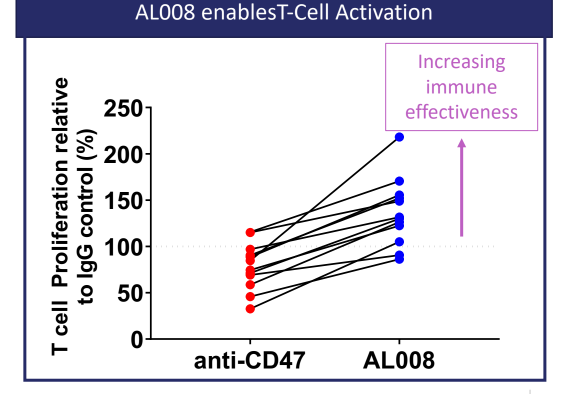
Re-acquiring rights granted to Innovent Biologics

PRODUCT CANDIDATE

- Selectively binds to multiple SIRP α variants
- Does not inhibit T-cell activator SIRPα

PRECLINICAL ACTIVITY

- Promotes T-cell proliferation more effectively than anti-CD47
- Demonstrates robust anti-tumor activity in multiple models (single agent and with checkpoint inhibitors)
- No decrease in red blood cells or platelets





Translating Scientific Insights into a Broad Portfolio of First-in-Class Programs

NOVEL APPROACH	Founded to pioneer a new field of research: Immuno-neurology	Informed by neuroscien human genetics and immunology	ce, Substantial IP portfolio established: 20 issued patents, 450+ patent applications
MULTIPLE CLINICAL TRIALS	Phase III Clinical Program for FTD-PGRN	Clinical Programs for AD, FTD-GRN, FTD- C9ORF72, ALS	Pre-Clinical Programs for AD, PD, Solid tumors
WORLD CLASS PARTNERS	\$700M upfront \$1.5B+ milestone 50-50 U.S. profit share Tiered double-digit royalties	\$20M \$986N	A upfront payment equity investment A milestone payments I 50-50 profit share
STRONG FINANCIALS	\$809 MILLION IN CASH		
alector ⁻	FTD = Frontotemporal dementia, PD = Parkinson's Disease, AD = Alzheimer's Disease, ALS = Amyotrophic lateral sclerosis		



Thank You