



Redefining Neuroscience Drug Development

August 2025

Important Disclosures

This presentation contains forward-looking statements about Neumora Therapeutics, Inc. (the “Company,” “we,” “us,” or “our”) within the meaning of the federal securities laws, including statements related to: Neumora’s intention to redefine neuroscience drug development by bringing forward the next generation of novel therapies that offer improved treatment outcomes and quality of life for patients; the timing, progress and plans for its therapeutic development programs, including the timing of clinical trial initiation and data readouts and upcoming milestones and catalysts; expectations and projections regarding future operating results and financial performance, including the sufficiency of its cash resources and expectation of the timing of its cash runway; and other statements identified by words such as “could,” “expects,” “intends,” “may,” “plans,” “potential,” “should,” “will,” “would,” or similar expressions and the negatives of those terms. Other than statements of historical facts, all statements contained in this press release are forward-looking statements within the meaning of the "safe harbor" provisions of the Private Securities Litigation Reform Act of 1995. These statements are subject to risks and uncertainties that could cause the actual results to be materially different from the information expressed or implied by these forward-looking statements, including, among others: the risks related to the inherent uncertainty of clinical drug development and unpredictability and lengthy process for obtaining regulatory approvals; risks related to the timely initiation and enrollment in our clinical trials; risks related to our reliance on third parties, including contract research organizations; risks related to serious or undesirable side effects of our therapeutic candidates; risks related to our ability to utilize and protect our intellectual property rights; and other matters that could affect sufficiency of capital resources to fund operations. For a detailed discussion of the risks and uncertainties that could cause actual results to differ from those expressed in these forward-looking statements, as well as risks relating to Neumora’s business in general, please refer to the risk factors identified in the Company’s filings with the Securities and Exchange Commission (SEC), including but not limited to its Quarterly Report on Form 10-Q for the quarter ended June 30, 2025 which was filed with the SEC on or about the date hereof. Forward-looking statements speak only as of the date hereof, and, except as required by law, Neumora undertakes no obligation to update or revise these forward-looking statements. Our results for the quarter ended June 30, 2025 are also not necessarily indicative of our operating results for any future periods.





Our Mission

We are focused on redefining neuroscience drug development by bringing forward the next generation of novel therapies that offer improved treatment outcomes and quality of life for patients suffering from brain diseases



Redefining Neuroscience Drug Development



**Industry leading
CNS pipeline with long-
dated IP into the 2040s**

**Multiple value-creating
clinical catalysts
on the horizon**



**Built at scale with strong
balance sheet; \$850M
raised since 2021**

**Cash runway into
2027 supporting
company growth**



**World-class team with
differentiated approach**

**Maximizing probability of
success with team and
proprietary approach**

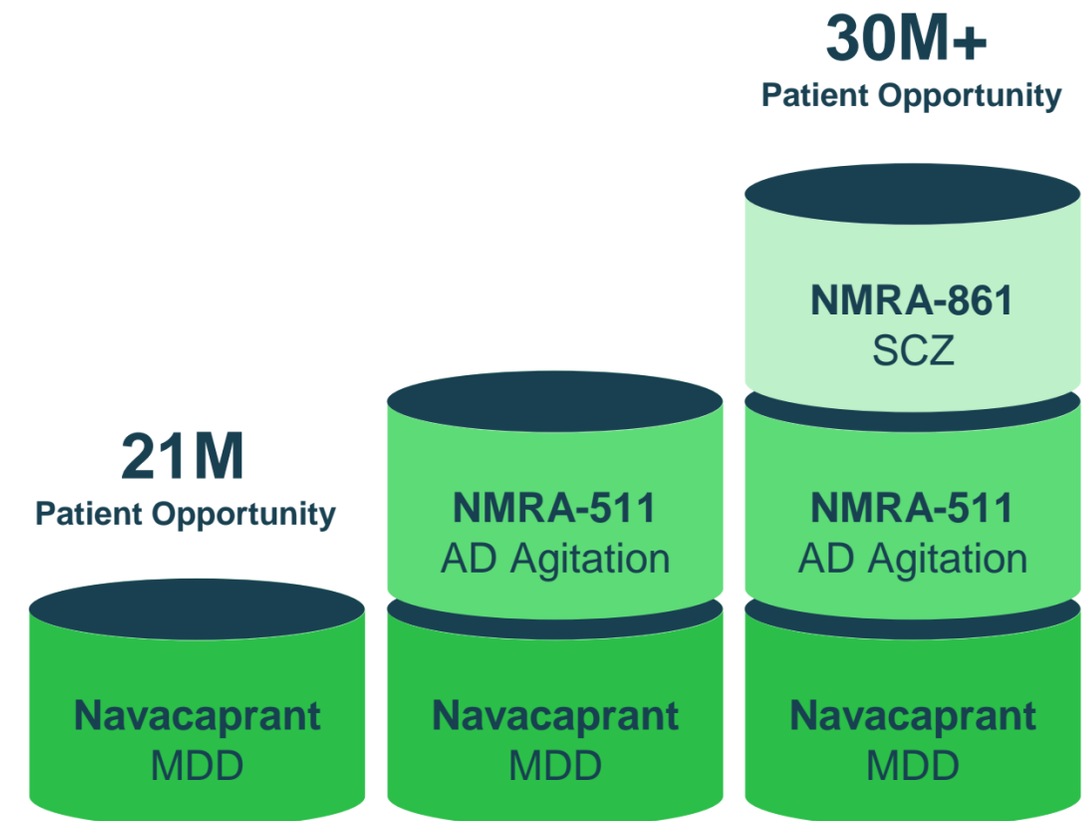
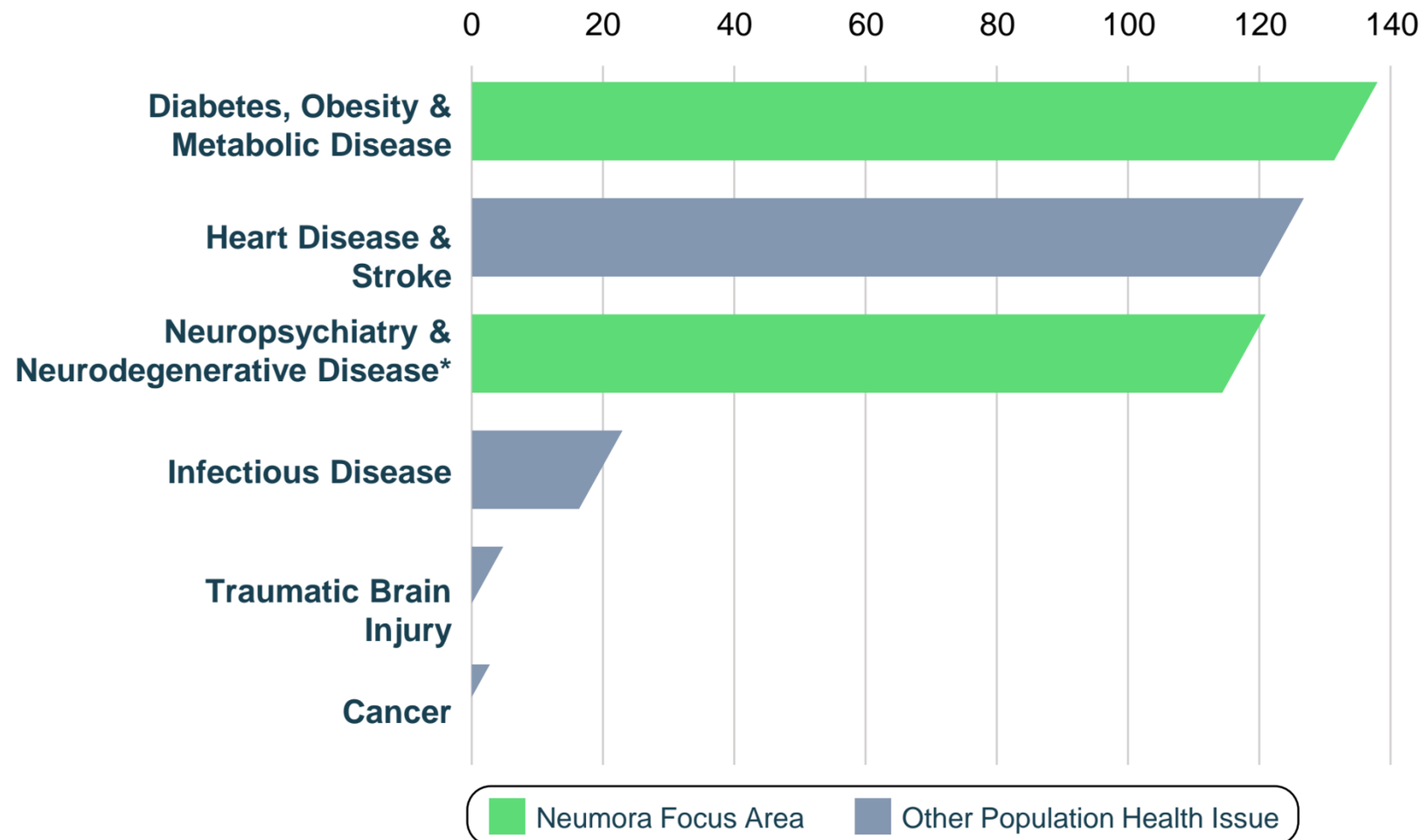


Neumora is Tackling Large Population Health Challenges

Neumora's clinical-stage pipeline has potential to reach up to ~30M+ patients with a robust IP runway into 2041+

Biggest Health Disorders Facing U.S.¹

Patients Impacted (M)



¹National Institutes of Health. Our Biggest Health Challenges. Accessed December 2023.

Note: Figure not intended as launch guidance or order. BPD = Bipolar Depression; MDD = major depressive disorder.

*Includes: MDD, BPD, Schizophrenia, Generalized Anxiety Disorder, Post Traumatic Stress Disorder, Substance Use Disorder, Alzheimer's Disease, Parkinson's Disease, Attention-Deficit Hyperactivity Disorder

Advancing a Leading Neuroscience Pipeline

- **Broad pipeline** addressing some of the most prevalent brain diseases
- Targeting novel mechanisms across a **broad range** of neuropsychiatric and neurodegenerative indications

PROGRAM <i>Target/Mechanism</i>	INDICATION <i>U.S. Prevalence</i>	Preclinical	Phase 1	Phase 2	Phase 3	MILESTONE <i>Guidance</i>
Navacaprant <i>KOR Antagonist</i>	Major Depressive Disorder 21M					KOASTAL-3, -2 topline data 1Q26, 2Q26
NMRA-511 <i>V1aR Antagonist</i>	Agitation in Alzheimer's Disease 6M					Phase 1b data around year-end 2025
NMRA-861 <i>M4 Modulator</i>	Schizophrenia 3M					Phase 1 SAD/MAD data 1Q26
NMRA-898 <i>M4 Modulator</i>	Schizophrenia 3M					Progress next M4 compound into the clinic 2025
NMRA-215 <i>NLRP3 Inhibitor</i>	Obesity/Parkinson's Disease 103M/1M					Report DIO data 2025 Initiate Phase 1 studies 1Q26
NMRA-GCASE <i>GCase Activator</i>	Parkinson's Disease 1M					
NMRA-CK1δ <i>CK1δ Inhibitor</i>	ALS/Parkinson's Disease 25K/1M					

ALS = Amyotrophic lateral sclerosis; CK1 δ = Casein Kinase I Isoform delta; GCase = Glucocerebrosidase; IP = Intellectual Property; KOR = kappa opioid receptor; M4 = Muscarinic Acetylcholine Receptor M4; NLRP3 = Nucleotide-binding Domain, Leucine-rich-containing Family, Pyrin Domain-containing-3; V1aR = Vasopressin 1a Receptor; DIO = diet induced obesity mouse model.
 **All dates are approximate / estimates / projections only

MDD Represents a Major Population Health Challenge

MDD is the leading cause of disability worldwide¹

280M

people worldwide have MDD¹

21M

adults in the U.S. have MDD²; the median onset is ~32.5 years of age

30 years

since a novel mechanism of action was approved for MDD

Many people have inadequate response to medication and experience tolerability issues

85%

of patients either don't receive pharmacological treatment or fail to achieve remission with first-line treatment³⁻⁷

70%

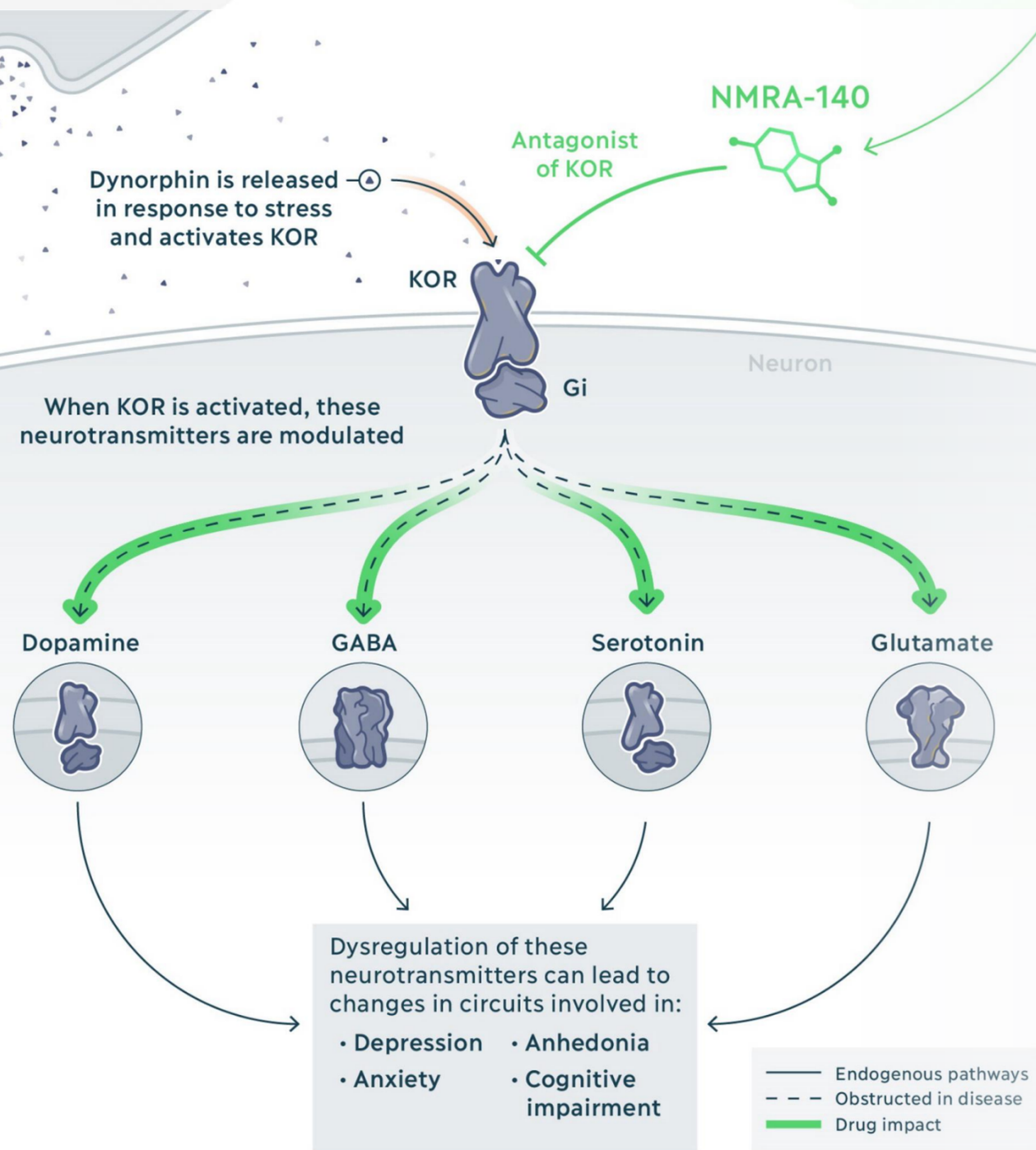
of people with MDD experience anhedonia⁸

60–85%

of patients treated with monotherapy⁹

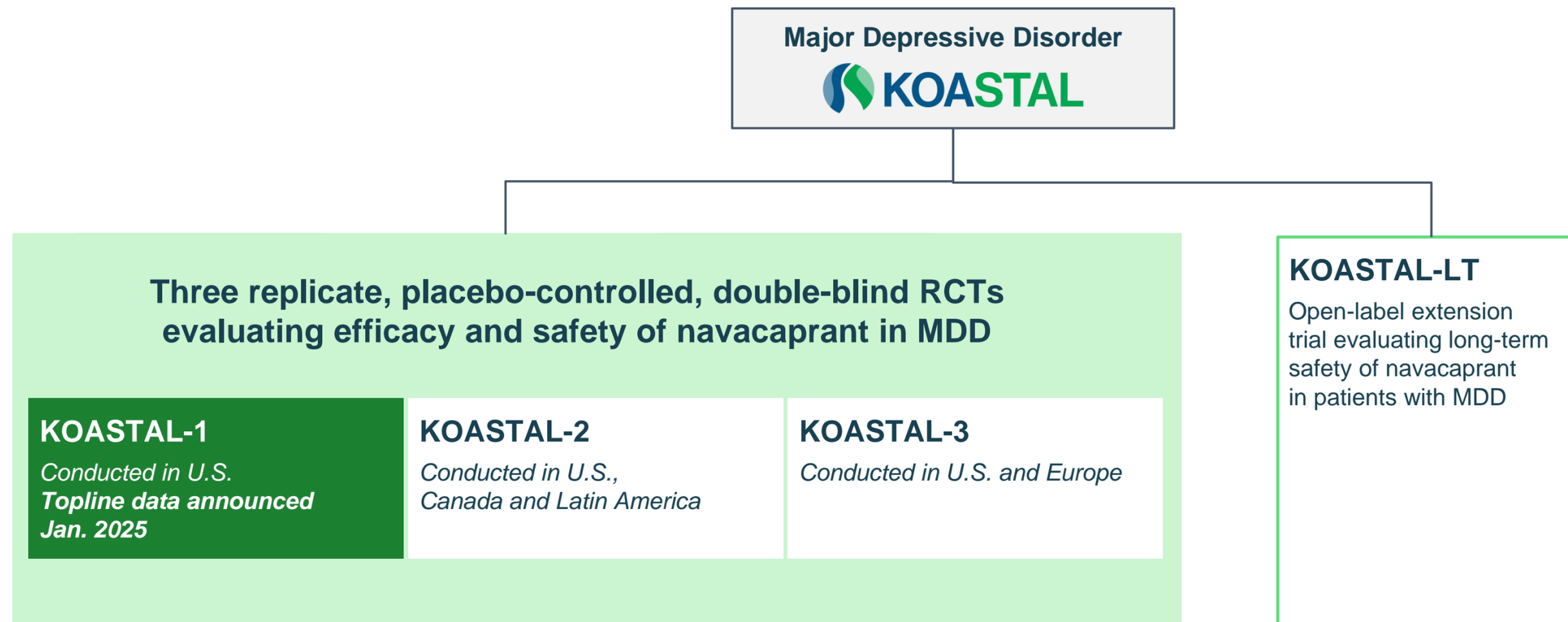


The Role of Kappa Opioid Receptor Antagonism in MDD



- The **kappa opioid receptor (KOR)** / dynorphin system is a well-characterized pathway, and results from preclinical studies support its potential to modulate depression, anhedonia, and anxiety
- KOR system overactivation in response to stress and mediation of depressive-like symptoms including anhedonia
- KOR antagonism may allow DA and 5HT release to return to adaptive levels during reward processing

Near-term Clinical Development Plan Focused on MDD with Opportunity for Further Expansion

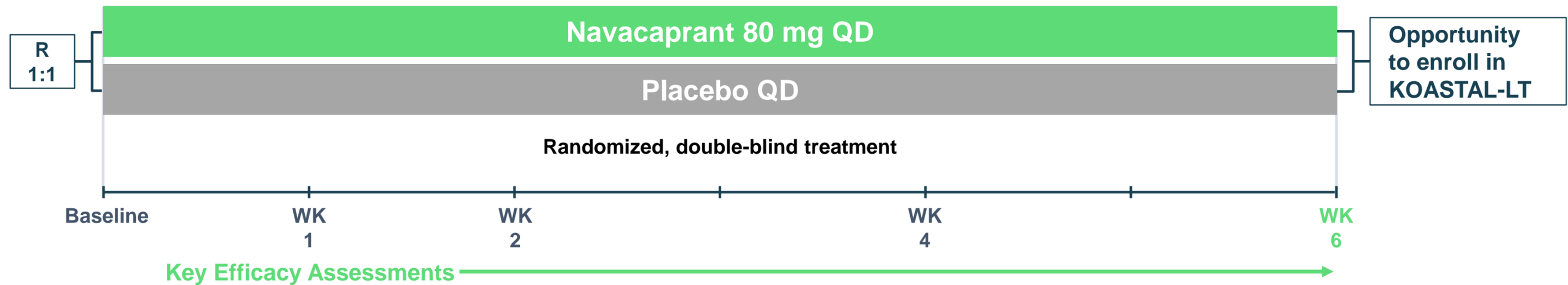


Additional indication opportunities include bipolar depression, substance use disorder, ADHD, Generalize Anxiety Disorder and Post-Traumatic Stress Disorder

KOASTAL Pivotal Study Design



KOASTAL Pivotal Efficacy Studies



KOASTAL-1, KOASTAL-2, KOASTAL-3 Summary

Inclusion Criteria:	<ul style="list-style-type: none"> Adults ages 18 – 65 diagnosed with MDD MADRS \geq 25 at baseline 	Other Secondary Endpoints Include: <ul style="list-style-type: none"> Δ from baseline to each timepoint in: <ul style="list-style-type: none"> CGI-S and CGI-I PHQ-9 HAM-A SDS
Primary Endpoint:	<ul style="list-style-type: none"> Δ from baseline to Week 6 in MADRS total score 	
Key Secondary Endpoint:	<ul style="list-style-type: none"> Δ from baseline to Week 6 in SHAPS total score 	Key Exploratory Endpoints*: <ul style="list-style-type: none"> Δ from baseline to each timepoint in: <ul style="list-style-type: none"> EQ-5D 5L WPAI-GH

*Safety Assessments include Change in Sexual Functioning Questionnaire (CSFQ-14)

Δ = Change; CGI-I = Clinical Global Impression-Improvement scale; CGI-S = Clinical Global Impression-Severity scale; EQ-5D 5L = EuroQol-5D 5L; HAM-A = Hamilton Anxiety Rating Scale; MADRS = Montgomery-Åsberg Depression Rating Scale; MDD = Major Depressive Disorder; PHQ-9 = Patient Health Questionnaire-9; QD = once daily; SDS = Sheehan Disability Scale; SHAPS = Snaith-Hamilton Pleasure Scale; wk = week; WPAI-GH = Work Productivity and Activity Impairment Questionnaire – General Health.

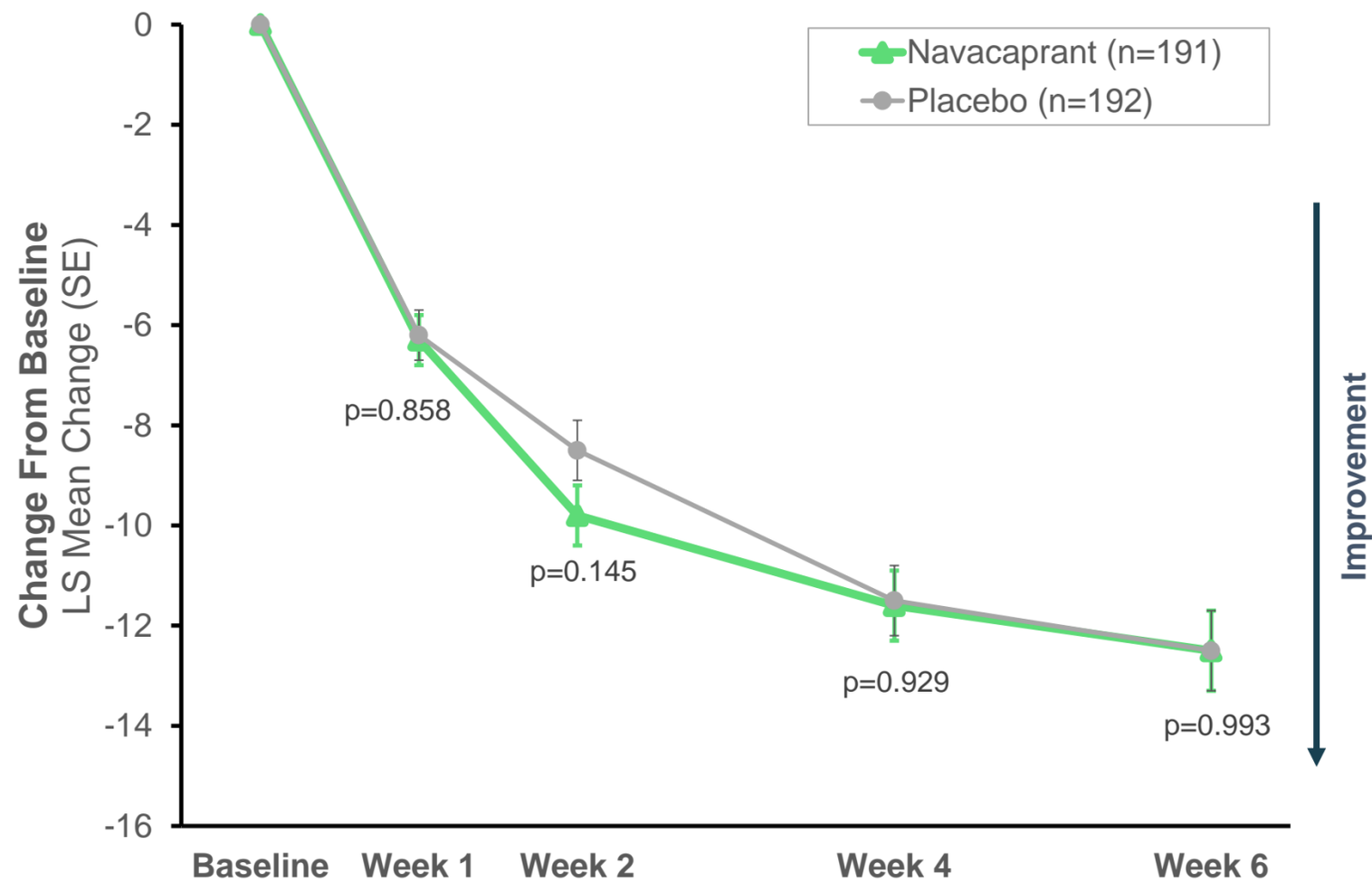
KOASTAL-1 Topline Data: Demographics and Baseline Characteristics

Intent-to-Treat Population	Navacaprant n = 191	Placebo N = 192
Age , mean (SD)	40.7 (14.0)	41.1 (13.2)
Sex , n (%)		
Male	86 (45.0%)	86 (44.8%)
Female	105 (55.0%)	106 (55.2%)
Race , n (%)		
White	112 (58.6%)	127 (66.1%)
Black or African American	38 (19.9%)	31 (16.1%)
Asian	25 (13.1%)	19 (9.9%)
Other	10 (5.2%)	10 (5.2%)
Missing/Unknown	6 (3.1%)	5 (2.6%)
Baseline MADRS total score , mean (SD)	32.2 (4.2)	32.8 (4.7)
Baseline SHAPS total score , mean (SD)	36.2 (6.2)	36.5 (6.7)

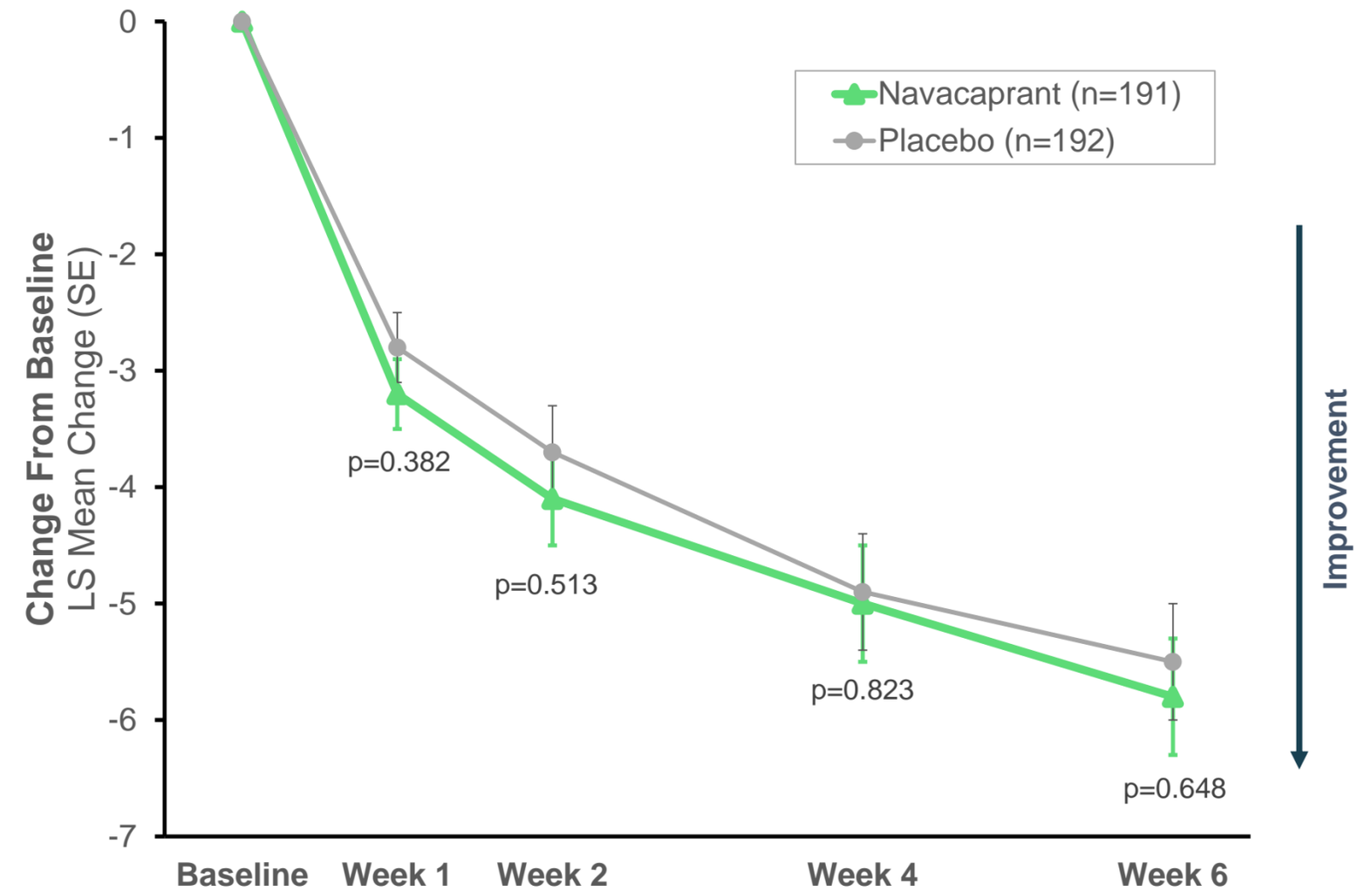


KOASTAL-1 Topline Data: Primary & Key Secondary Endpoint

MADRS Total Score
Intent-to-Treat Population



SHAPS Total Score
Intent-to-Treat Population



KOASTAL-1 Topline Data: Favorable Safety Profile Demonstrated

Navacaprant was safe and generally well tolerated, with no serious adverse events reported

TEAEs Incidence (>2% in either treatment group)	Placebo n=192	Navacaprant n=191
Preferred Terms	n (%)	n (%)
Headache	14 (7.3%)	13 (6.8%)
Diarrhea	4 (2.1%)	10 (5.2%)
Nasopharyngitis	8 (4.2%)	7 (3.7%)
Pruritus	4 (2.1%)	7 (3.7%)
Nausea	6 (3.1%)	6 (3.1%)
Constipation	6 (3.1%)	5 (2.6%)
Insomnia	4 (2.1%)	3 (1.6%)
Fatigue	9 (4.7%)	2 (1.0%)
Upper respiratory tract infection	6 (3.1%)	2 (1.0%)
Dizziness	5 (2.6%)	2 (1.0%)
Dry mouth	4 (2.1%)	2 (1.0%)
Somnolence	4 (2.1%)	2 (1.0%)
Urinary tract infection	4 (2.1%)	2 (1.0%)
Back pain	5 (2.6%)	0

- No signal for increased suicidal ideation or suicidal behavior¹
- Low discontinuation rate due to TEAEs (navacaprant 2.1%; placebo 3.1%)
- 83.3% of navacaprant-treated patients who completed 6 weeks' treatment elected to enroll in KOASTAL-LT



1. As measured by Columbia Suicide Severity Rating Scale (C-SSRS)

Optimizing KOASTAL-2 and -3 Phase 3 Trials

Topline data from KOASTAL-3 in the first quarter of 2026 and -2 in the second quarter of 2026



Site Selection

Adjusted clinical sites included in studies, with goal of including sites with demonstrated expertise in conducting MDD studies



Medical Monitoring

Using clinician-rated Massachusetts General Hospital Clinical Trials Network and Institute SAFER approach to verify the diagnosis and appropriateness of patient population



Screening Tools

Verified Clinical Trial (VCT) screening database complements the Clinical Trial Subject (CTS) database to screen for people who participate in multiple clinical trials



Target Enrollment

Option included in KOASTAL-2 and -3 protocols to overenroll the studies up to 25%



NMRA-511 is a Best-in-Class Vasopressin 1a Receptor Antagonist with Broad Potential Across Neuropsychiatric Disorders

Rationale

Vasopressin plays a role in the regulation of aggression, affiliation, stress and anxiety response

Indication

Agitation in Alzheimer's disease

Status

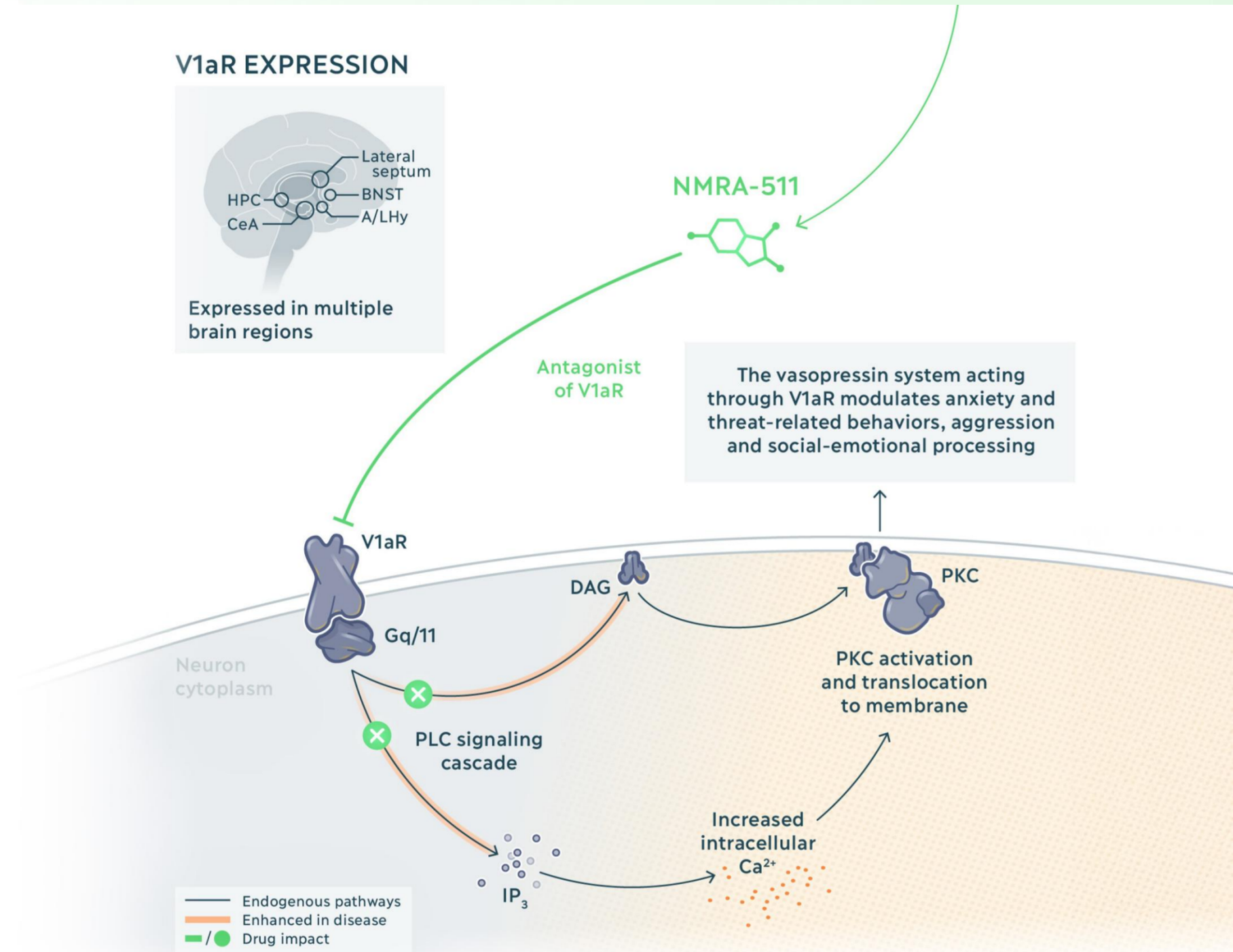
Phase 1b study underway with data anticipated around the end of 2025

Drug Profile

Oral, BID dosing

Strong IP Protection

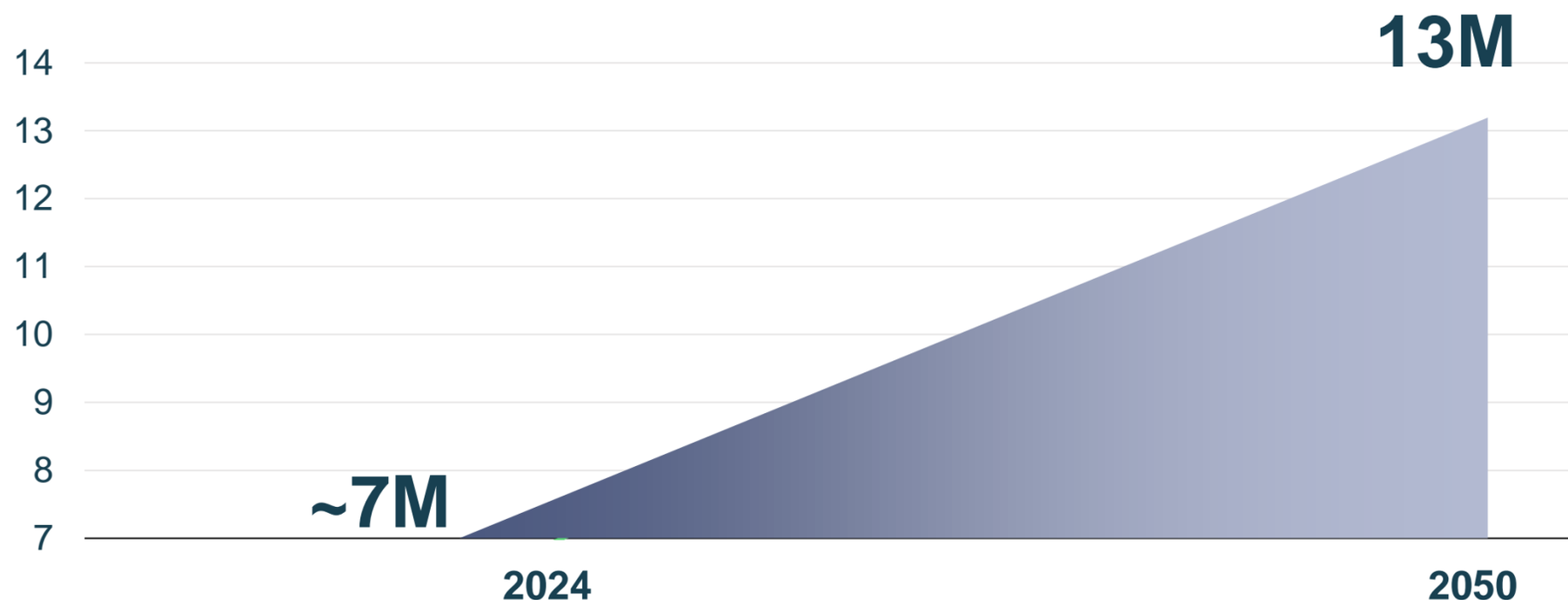
Expect exclusivity through 2042+, based on composition of matter protection and estimated patent term extension



Alzheimer's Disease Agitation Represents Large Market Opportunity with Significant Unmet Need

Agitation in Alzheimer's disease impacts a significant portion of the U.S. population; that number is expected to increase as the population ages¹

U.S. Adults with Alzheimer's Disease (M)¹



>70%

of people with AD experience agitation at some point in their disease²

Significant unmet medical need exists in this population³

Agitation is among the most disruptive symptoms of AD. It is associated with greater caregiver stress, increased morbidity and mortality and earlier placement in long-term care facilities. The only currently approved product carries a black-box warning for mortality in elderly people.

¹Alzheimer's Association. Alzheimer's Disease Facts and Figures. May 2024. ²Ijaopo et al., 2017., Translational Psychiatry.; ³Koenig et al., 2016, Current Psychiatry.

Several Lines of Evidence Indicate that V1a Receptor Antagonists Have Therapeutic Potential for Reducing Symptoms of Agitation



The vasopressin system modulates social-emotional, anxiety and threat-related behaviors across species

- V1aR expression patterns critically affect social behavior¹⁻⁵
- Rodent selection lines bred for aggression or anxiety show dysregulated vasopressin release and HPA axis functioning⁶
- Vasopressin-deficient rodents display impaired responses to threat stimuli, reduced anxiety and depressive-like behaviors, and impaired aggression toward intruders⁷⁻⁹



In healthy volunteers, vasopressin enhances reactivity to threatening stimuli and disrupts emotional control¹⁻²

- Exogenously administered vasopressin increases autonomic responsiveness to threat stimuli and increases anxiety²
- V1a antagonist administration suppresses anxiety induced by unpredictable threats¹⁰



Positive association between vasopressin and aggression in people with personality disorders¹¹

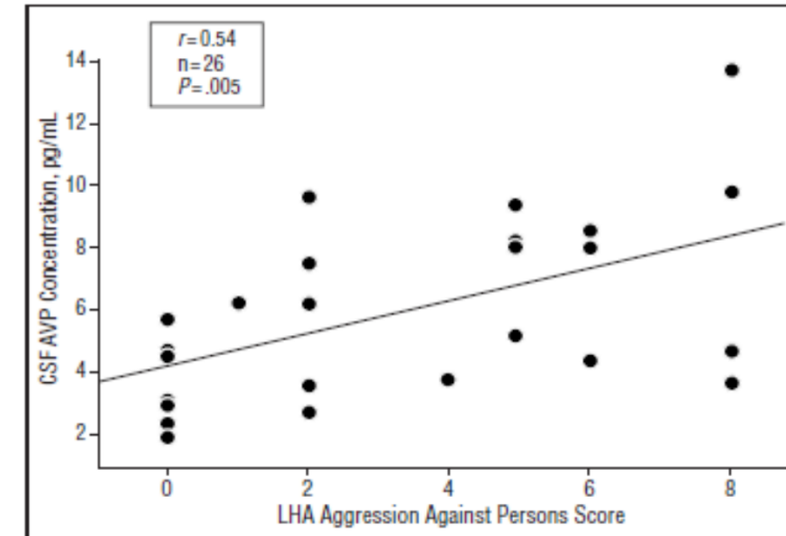


Figure 1. Correlation between Aggression Against Persons (the fighting and assault items) scores on the Life History of Aggression (LHA) assessment and cerebrospinal fluid (CSF) arginine vasopressin (AVP) concentrations in 26 individuals who met the DSM-IV criteria for personality disorder.



In HD irritability, an investigational V1a receptor antagonist reduced an exploratory endpoint measuring aggression¹²

Together, these data support the development of a V1a receptor antagonist for the treatment of symptoms of agitation, aggression, and anxiety

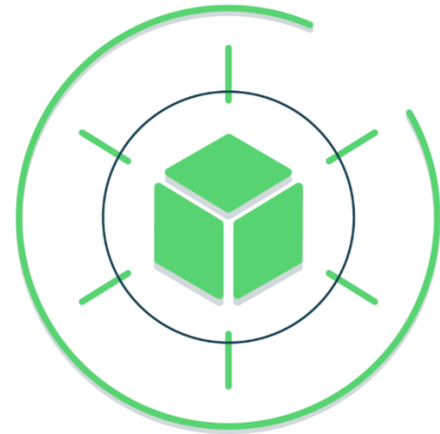
¹Ebstein et al., 2009, New York Academy of Sciences.; ²Thompson et al., 2006, PNAS.; ³Insel et al., 2010, *Neuron Review*, PNAS; ⁴Carter et al., 1995, *Neuroscience Biobehavioral Review*.; ⁵Wang et al., 1994, PNAS.; ⁶Veenema and Neumann, 2007, *Brain behavior, evolution*.; ⁷Zelena et al., 2009, *Journal of Endocrinology*.; ⁸Mlynarik et al., 2007, *Hormones and Behavior*.; ⁹Fodor et al., 2014, *Psychoneuroendocrine*.; ¹⁰Lago et al., 2021, *Psychopharmacology*.; ¹¹Coccaro et al., 1998., *JAMA Psychiatry*.; ¹²Maibach et al., 2022, *Personalized Medicine*.
HPA = hypothalamic-pituitary-adrenal

NMRA-511 Profile Supports Advancement into Alzheimer's Disease Agitation



Best-in-Class Pharmacology¹

- Highly potent at V1a
- High selectivity over V1b, V2, and oxytocin receptors
- Excellent brain penetration



Strong Pre-Clinical Data²

- Robust pharmacodynamic (PD) effect in rodents
- Robust activity in a marmoset 'human threat test' model of stress/anxiety



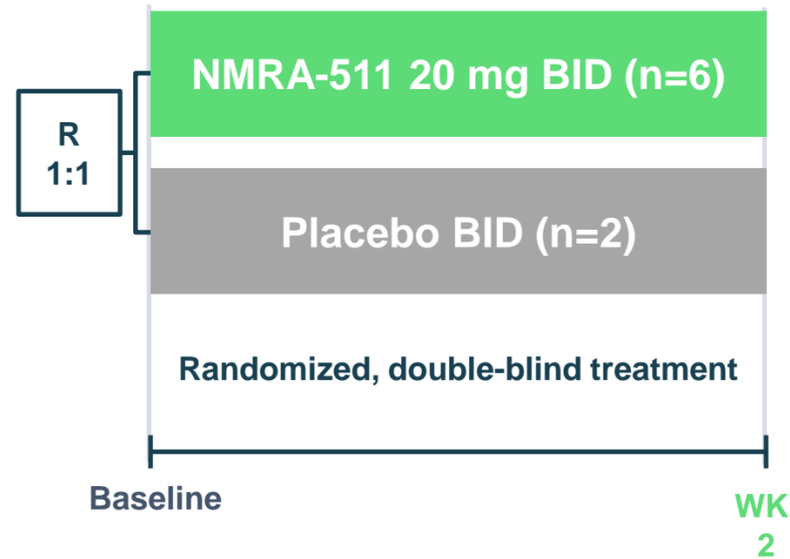
PK and Safety Data from Phase 1 Support Advancement¹

- NMRA-511 was safe and very well-tolerated in Phase 1 SAD/MAD study
- NMRA-511 was safe and well-tolerated in healthy elderly volunteers

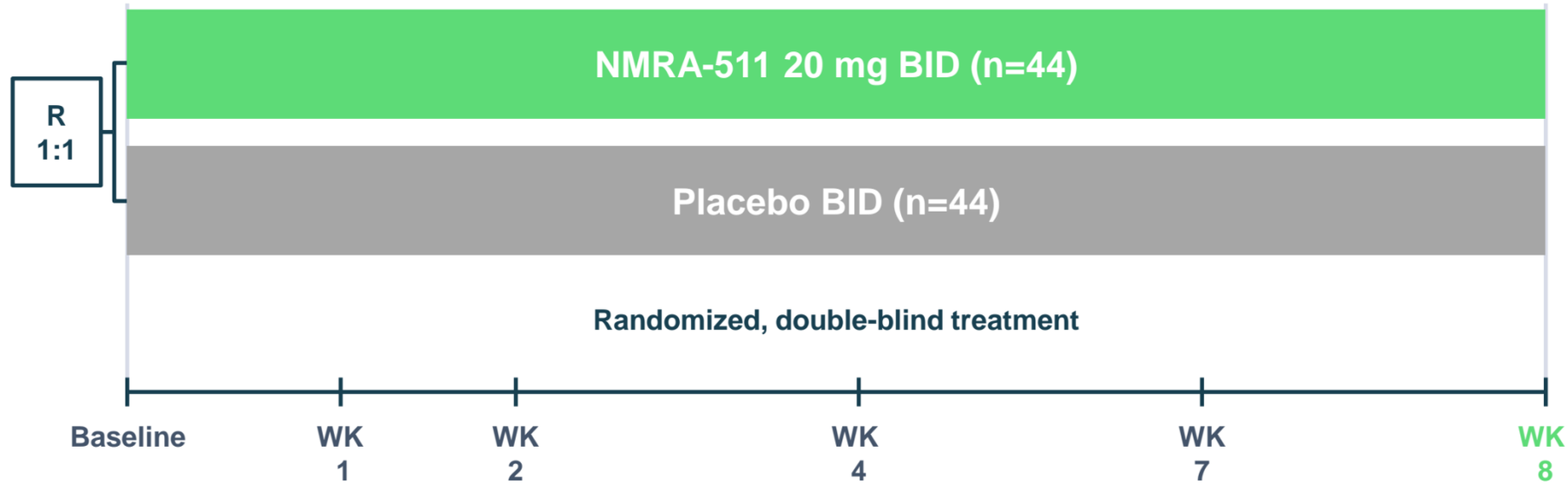


NMRA-511 Signal Seeking Study in Alzheimer's Disease Agitation

Part A: 2-Week Evaluation Period Enrolling Healthy Elderly Participants



Part B: 8-Week Evaluation Period Enrolling People with Alzheimer's Disease Agitation (ADA)



NMRA-511 Phase 1b Study

- | | |
|---|---|
| Part A Inclusion Criteria: | <ul style="list-style-type: none"> • Healthy elderly adult participants aged 65-80 years |
| Part B Inclusion Criteria: | <ul style="list-style-type: none"> • Adults aged 55-90 years with mild-severe dementia (MMSE score of 5-24) and clinically significant agitation (CMAI total score 45-100) |
| Part B Primary Endpoint: | <ul style="list-style-type: none"> • Δ from baseline to Week 8 in CMAI total score |
| Part B Other Endpoints Include*: | <ul style="list-style-type: none"> Δ from baseline to Week 8 in: <ul style="list-style-type: none"> • CGI-S Agitation total score • mADCS-CGIC total score • Caregiver Diary of participant agitation, aggression, and/or anxious behaviors • NPI total score |
| Statistics: | <ul style="list-style-type: none"> • Study not powered to demonstrate statistical significance • Designed as a signal-seeking study; effect size will inform the potential future development of NMRA-511 in ADA |

*Safety Assessments include adverse events, clinical laboratory, vital signs, physical examination, 12-lead electrocardiogram (ECG), Columbia-Suicide Severity Rating Scale (C-SSRS).
 Δ = Change; BID = twice daily; CMAI = Cohen-Mansfield Agitation Inventory; MMSE = Mini-Mental State Examinations; CGI = Clinical Global Impression of Change for Agitation; mADCS-CGIC = mADCS-CGIC Agitation modified Alzheimer's Disease Cooperative Study – Clinical Global Impression of Change for Agitation; NPI = Neuropsychiatric Inventory.

M4 PAM Franchise: Differentiated M4R PAMs for Schizophrenia

NMRA-861 Target Profile

Pharmacology

Neumora has multiple series of chemically distinct, highly selective M4 muscarinic receptor PAMs, including NMRA-861, designed for antipsychotic-like efficacy with the potential for improved tolerability profile

Indication

Schizophrenia

Target Administration

Oral, once-daily

IP

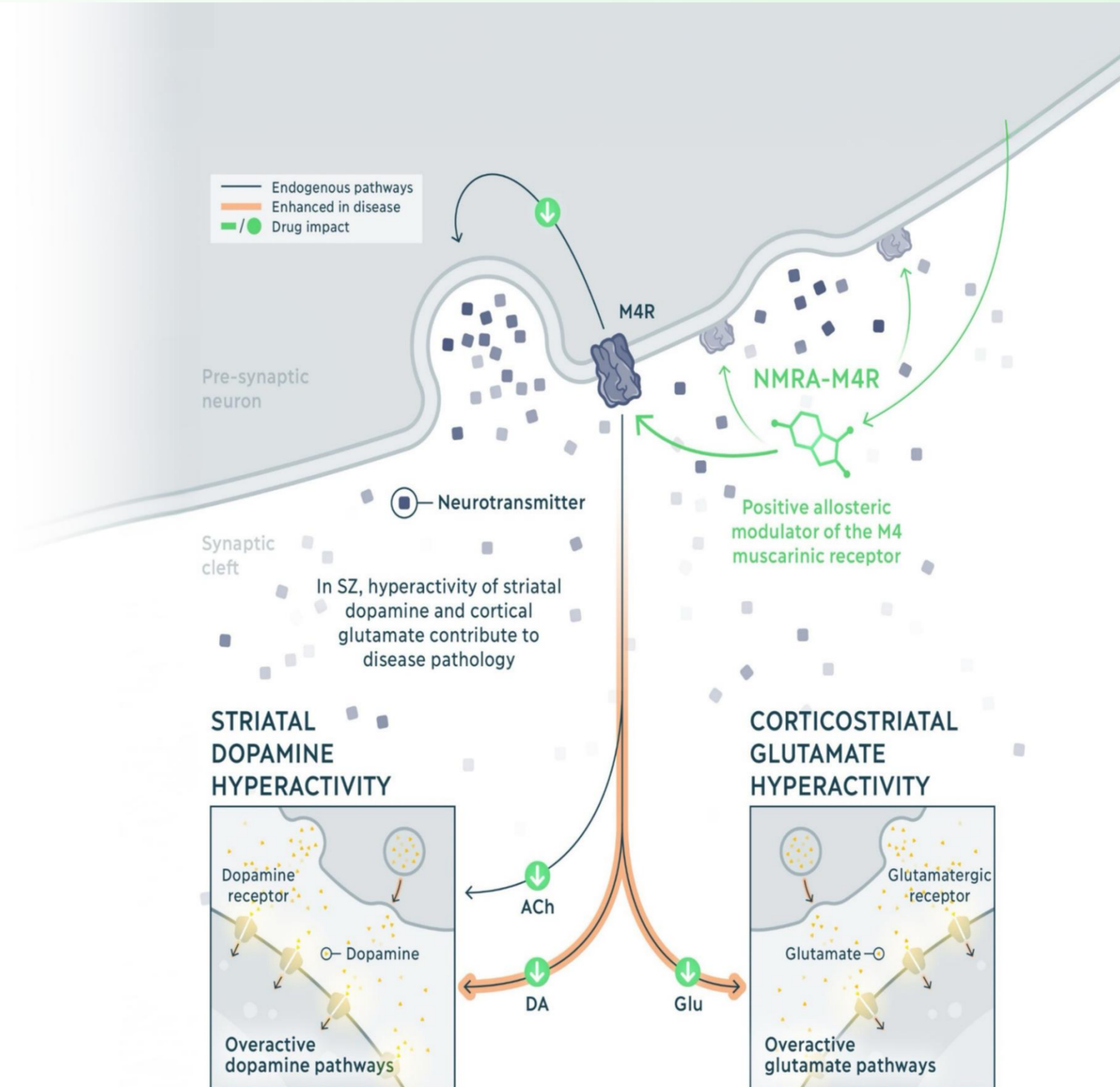
Composition of matter patent extending to 2044+*

Epidemiology

Estimated 3 million patients in the U.S. with schizophrenia¹

Expected Milestones

Report Phase 1 SAD/MAD data for NMRA-861 in the first quarter of 2026



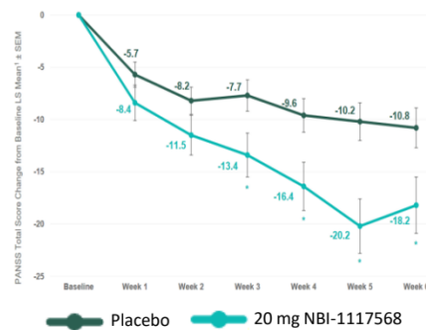
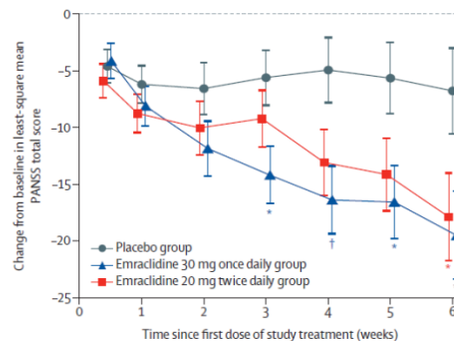
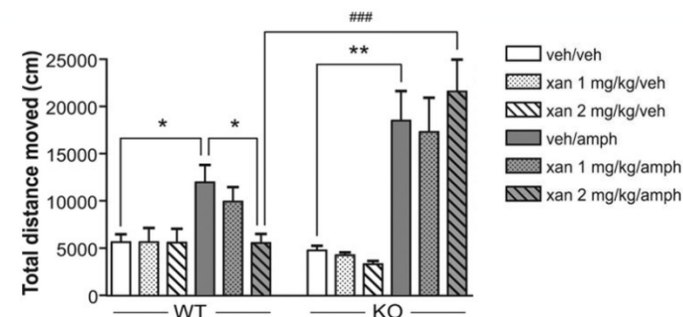
¹Wander, C. *Am J Manag Care*. 2020;26:S62-S68.

*Excluding any patent term adjustment or extension
PAM = positive allosteric modulator

An Optimized Muscarinic Drug Profile Would Include Selectivity and Potency in the CNS

Preclinical data and clinical data in acute schizophrenia supports M4 as a driver of antipsychotic activity

Activity of xanomeline (active component of Cobenfy™) is dependent on M4R in mice



Non-selective muscarinic agents are associated with a range of peripheral AEs

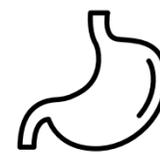
M4



Cardiovascular

Transient increased BP & heart rate

M1, M2, M3



GI Tract

Increased gastric secretion & gastric motility

M1, M2, M3



Cardiovascular

Direct effect on cardiac function – increased BP & heart rate

M1, M3



Glands

Increased salivation
Increased lacrimation
Increased sweating

PAMs offer the benefits of greater selectivity

- Targeting the allosteric site specifically allows for greater selectivity for M4 over other muscarinic sub-types than if targeting the orthosteric site due to binding site conservation
- To date the pharmacology of agonists targeting the orthosteric site are often thought to display 'partial' agonism which could contribute to variable clinical responses
- PAMs allow for more precise potentiation of M4, maintaining the spatial and temporal signaling dynamics of ACh

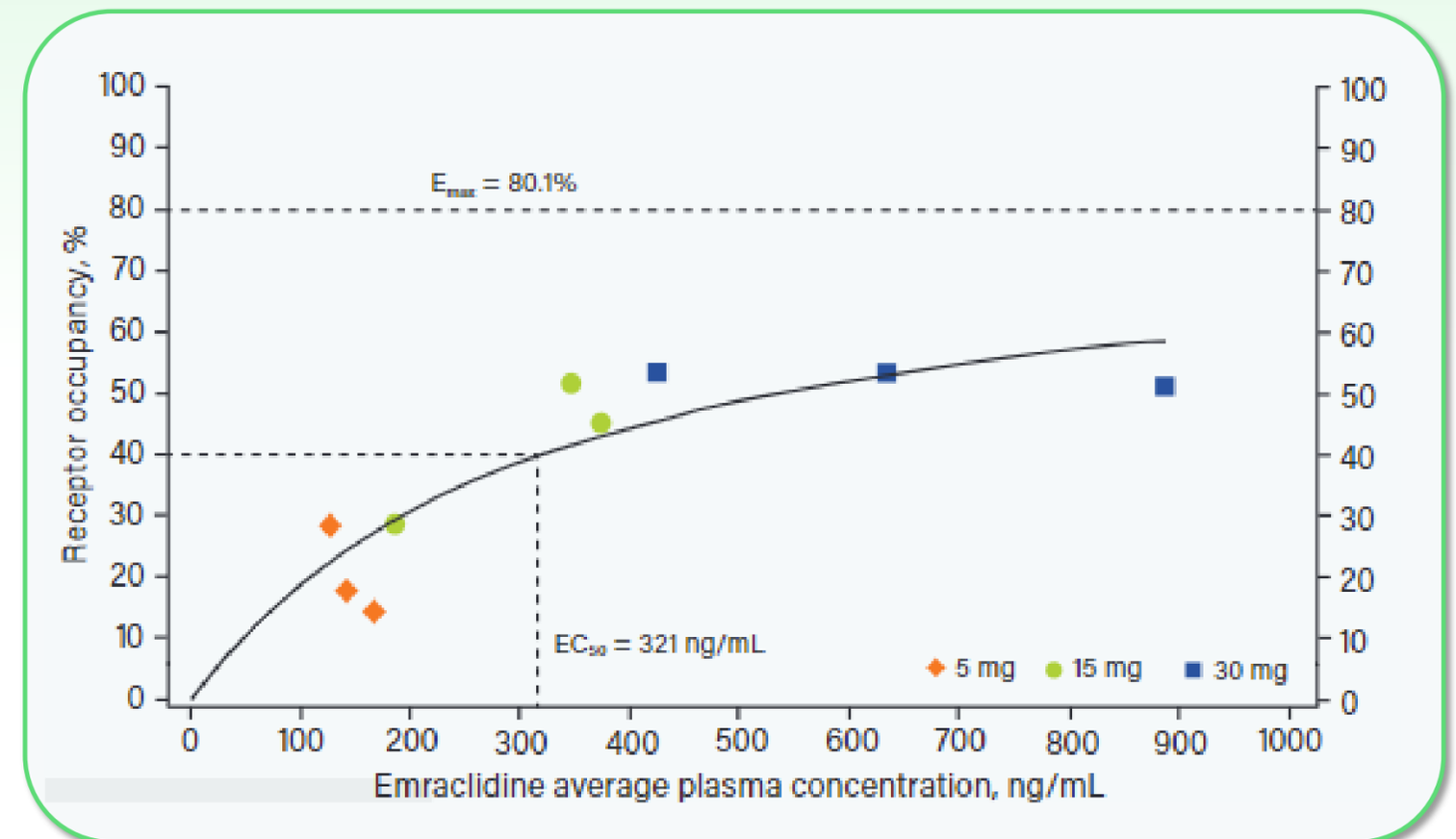


Emraclidine Receptor Occupancy Disconnected from Plasma Exposures

Receptor Occupancy for Emraclidine in Humans Suggests the Compound has Limited Brain Exposure

- In a human PET study, peripheral concentration of emraclidine shows dose linear response
- However, CNS receptor occupancy unchanged when dose doubled from 15 to 30 mg
- Data suggests emraclidine may have limitations in engaging the M4 receptor in the brain

Low CNS exposure could limit efficacy



CNS = central nervous system; PET = positron emission tomography

Duvvuri S, et al. *Evaluation of M4 Muscarinic Receptor Occupancy by Emraclidine Using [¹¹C]MK-6884 PET in Healthy Volunteers*. Poster M206. Presented at the 62nd Annual Meeting of the American College of Neuropsychopharmacology, Tampa, FL: December 3 – 6, 2023.

NMRA-861 Has Potential Best-in-Class Potency and Optimized Brain Penetration

NMRA-861 and –898 potentially more potent than emraclidine across multiple assays

NMRA-861 and –898 are selective for M4 over other muscarinic receptor subtypes

Neumora M4 PAMs are optimized for high CNS exposure

Neumora M4 PAMs are optimized for once daily dosing

Convulsions have not been observed with NMRA-861 or –898

		NMRA-861 ¹	NMRA-898 ¹	Emraclidine
	M4 EC ₅₀ (human; cAMP) ¹	6 nM	13 nM	26 nM
	M4 EC ₅₀ (human; Ca ²⁺) ¹	2 nM	8 nM	180 nM
	Selectivity at other muscarinic receptor subtypes (EC ₅₀) ¹	M1, M3, M5 > 10 μM, M2 0.7 μM	M1, M2, M3, M5 > 10 μM	M1, M3, M5 > 10 μM, M2 5.7 μM
	Brain exposure MDCK permeability (target >10) P-gp efflux ratio (target <2) ^{1,2}	High 45.5 1.26	High 36.7 0.93	Moderate 9.5 3, 6.02 ^{1,2}
	Human half-life ³	Pending Phase 1 Study	Pending Phase 1 Study	9 – 12 hr
	Preclinical convulsions	Not observed in rat, dog or rabbit	Not observed in rat, dog or rabbit	Unknown

NMRA-861 has potential best-in-class pharmacology and clinical differentiation

Note: Data on this slide is presented for illustrative purposes only. These molecules have not been studied in head-to-head clinical trials.

cAMP = cyclic adenosine monophosphate; CNS = central nervous system; PAM = positive allosteric modulator

1. Data generated by The Warren Center for Neuroscience Drug Discovery at Vanderbilt University on behalf of Neumora across NMRA-861, NMRA-898 and emraclidine. 2. Butler CR, et al. *J Med Chem.* 2024 Jul 11;67(13):10831-47. 3. Krystal JH, et al. *Lancet.* 2022 Dec 17;400(10369):2210-20.

SAD/MAD Evaluating NMRA-861 in Healthy Adults and People with Stable Schizophrenia

Study Objectives

- Confirm once-daily dosing – based on PK profile in humans
- Evaluate tolerable doses in people with stable schizophrenia
- Establish CNS penetration – based on CSF exposure

SAD – Part 1 CSP

	Dose Cohorts ^a	Participants	Randomization
Part 1A	5 mg, 15 mg, 45mg, 80 mg, final dose TBD ^b	Healthy adults	6:2 active:placebo
Part 1B (Fed-Fasted cohort)	Dose to be determined	Healthy adults	12 active

MAD – Part 2 CSP

	Dose ^c	Participants	Randomization
Cohort 1	15 mg QD	Healthy adults	6:2 active:placebo
Cohort 2	45 mg QD	Healthy adults	
Cohort 3	Dose to be determined	Healthy adults OR with stable schizophrenia ^d	
Cohort 4	Dose to be determined	Healthy adults OR with stable schizophrenia ^d	
Cohort 5	Dose to be determined	Adults with stable schizophrenia	

■ Healthy adults
 ■ Adults with stable schizophrenia

^aSentinel dosing done in each cohort. ^bAdditional optional cohort dose to be decided. ^cSentinal dosing only if recommended by SRC. ^dParticipants enrolled into flexible cohorts will be determined by SRC recommendation.

NMRA-215: Differentiated NLRP3 Inhibitor for Obesity and Related Metabolic Diseases

NMRA-215 Target Profile

Rationale/Pharmacology

NLRP3-related inflammatory response via release of IL-1 β , IL-18 and IL-6 cytokines is associated with obesity^{1,2}

Indication

Obesity, Parkinson's Disease

Target Administration

Oral, once-daily

IP

Composition of matter patent extending to 2043+*

Epidemiology

~1.13 billion patients in the world with obesity by 2030³

Expected Milestones

- Report preclinical data in 2025
- Progress NMRA-215 into the clinic in 1Q 2026

Multiple factors drive NLRP3-mediated inflammation resulting in disease

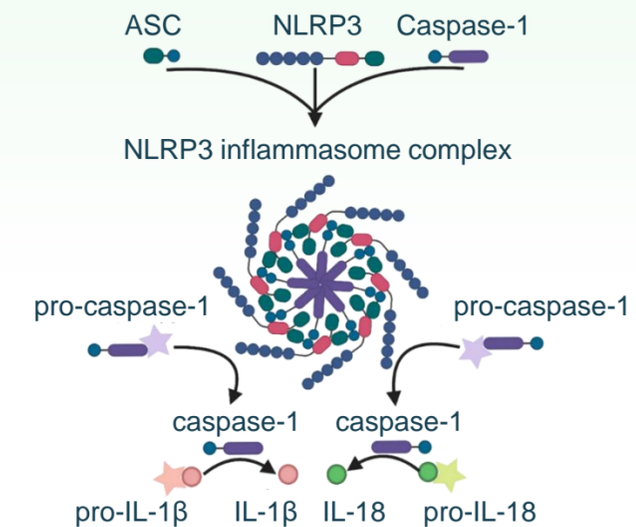


DRIVERS



- Diet (e.g., lipids)
- Environment
- Genetics
- Aging

NLRP3 Activation



DISEASES



- Neurodegeneration (Parkinson's)
- Cardio-metabolic (obesity)
- Monogenic / autoimmune (CAPS)

*Excluding any patent term adjustment or extension

1. O'Brian et al. *J Neuroinflammation*. 2020;17(1):104. 2. Wani K et al. *Int J Environ Res Public Health*. 2021;18(2):511. 3. World Obesity Federation. World Obesity Atlas 2024. London: World Obesity Federation, 2024. <https://data.worldobesity.org/publications/?cat=22>.

2. AdipoGen Life Sciences. <https://adipogen.com/inflammasomes/rce>



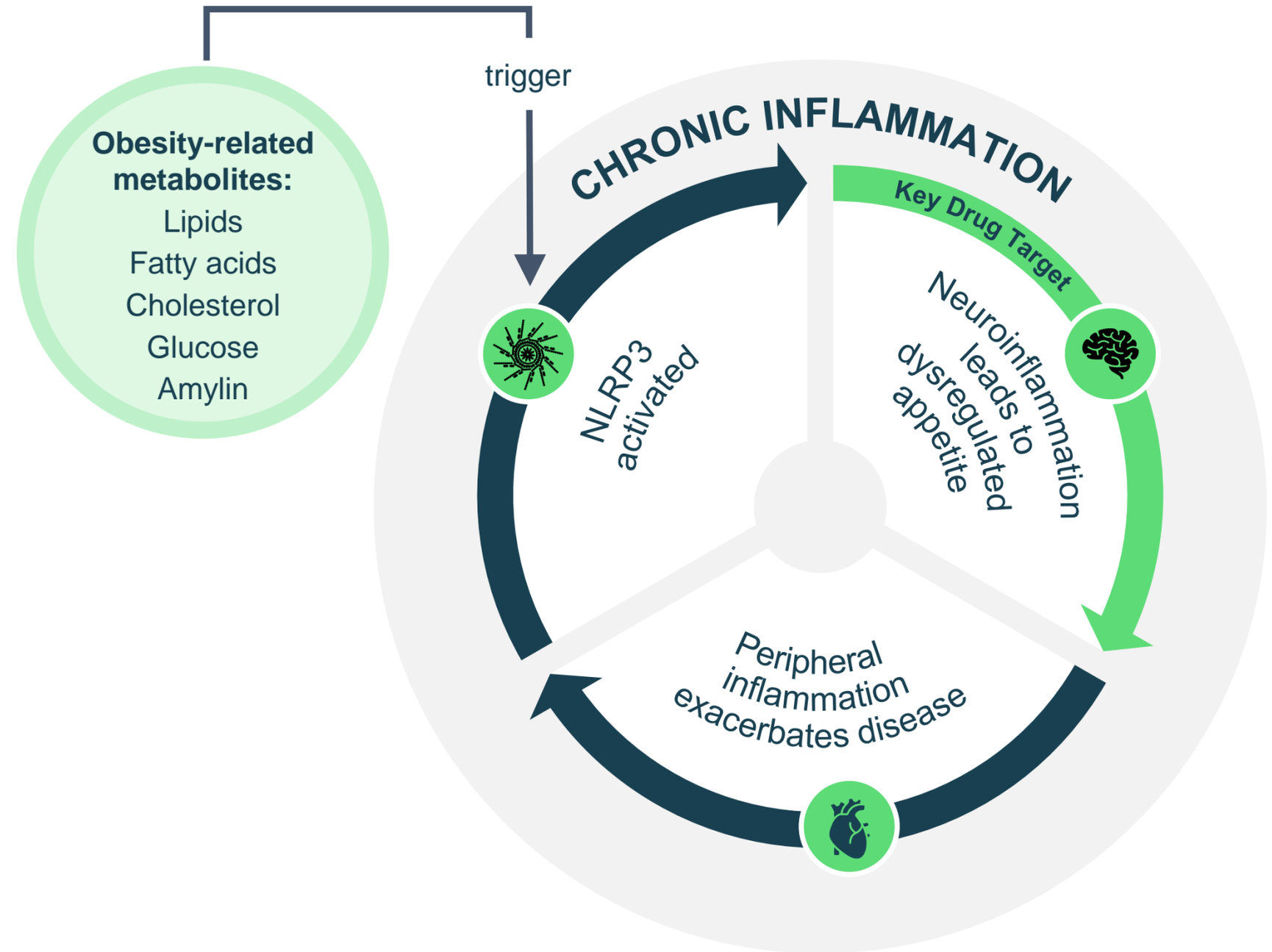
Obesity is Driven by Inflammation and Represents One of the Greatest Public Health Challenges



By 2030,
1.13 BILLION
people worldwide will be living with obesity¹



Driving a significant market
for obesity treatments
\$130 - \$170 BILLION
estimated obesity market size in 2030



1. World Obesity Federation. World Obesity Atlas 2025. London: World Obesity Federation, 2025. <https://data.worldobesity.org/publications/world-obesity-atlas-2025-v7.pdf>

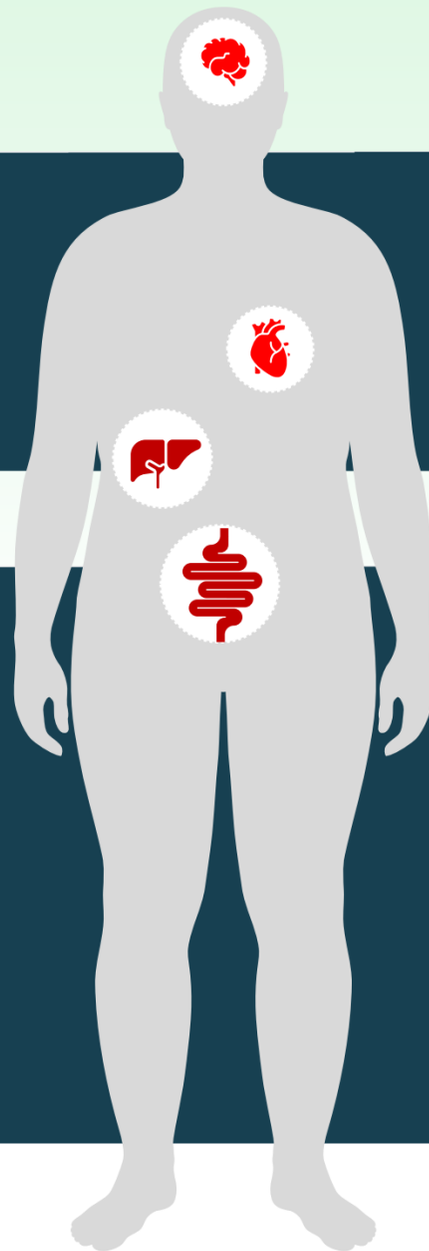


CNS Penetrant NLRP3 Inhibition Provides Systemic Benefit

System

CNS

Periphery



Drug Impact

Reduce neuroinflammation in the brain

Protect organ and vascular system from inflammation-related damage

Outcome

Reduced appetite and drive body weight loss

Reduce the risk of co-morbid diseases. NLRP3 inhibition:

- Reduces heart disease: improved CV outcomes in a clinical study
- Improves type II diabetes: reduced insulin resistance in mice

Potential treatment benefits driven by NLRP3 inhibition and positive effects of weight loss



NMRA-215 Has an Optimized Pharmacological Profile Including Best-in-Class CNS Exposure

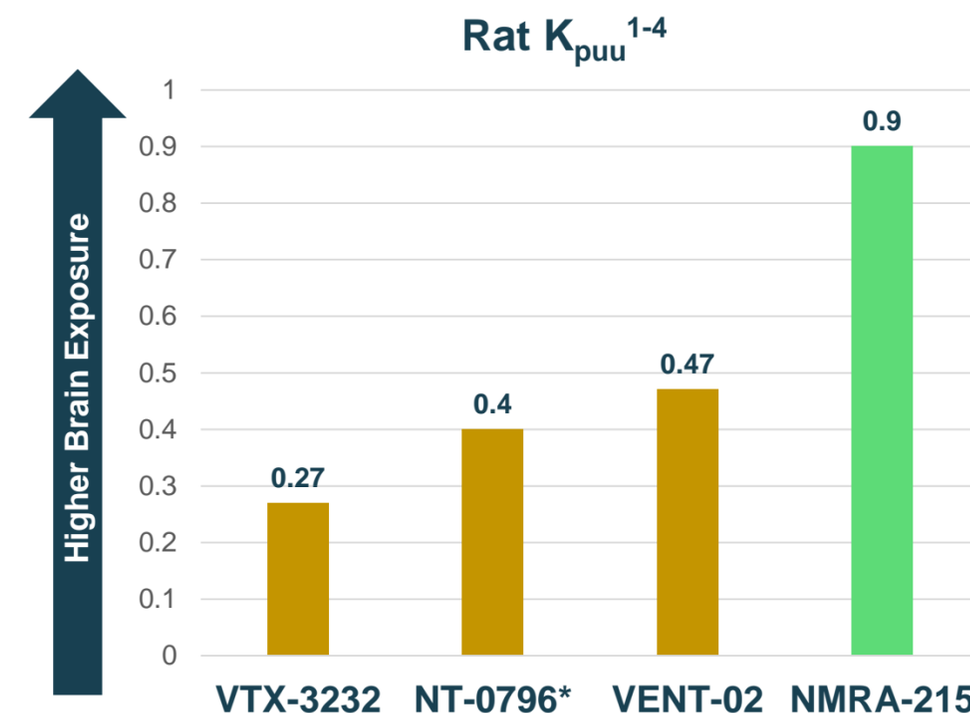
NMRA-215 is highly potent with low nM potency across a range of assays

NMRA-215 Assay Format	IC ₅₀
iMicroglia (IL-1β)	8 nM
THP-1 (IL-1β)	3 nM
Human whole blood (IL-1β)	16 nM
Target engagement (Nanobret)	5 nM

NMRA-215 is highly selective for NLRP3

- NMRA-215 is highly selective for NLRP3 versus other inflammasomes (NLRP1, NLRC4, AIM2)
- >250-fold selective for NLRP3 versus a broad panel of targets (Eurofins SafetyScreen87)
- Clean profile in cardiac ion channel and kinase screening panels

NMRA-215 is extensively characterized and optimized for brain exposure



MDCK permeability:	Unknown	14.0
P-gp efflux ratio:	Unknown	1.1

*NT0796 = mouse K_{puu}

1. Neumora data on file. 2. Thornton P, et al. *JPET*. 2024 Feb 15;388(3):813-826. 3. Ventus Data Presented at 5th Annual Inflammasome Summit. November 28 – 30, 2023. Boston, MA. 4. Ventyx R&D Day Presentation. Published Jan 2023.

Study Designs Maximize Hypothesis Testing in the Mouse DIO Model

Neumora studies will evaluate 3 hypotheses:

	Hypothesis	Value proposition
1	NMRA-215 as weight loss monotherapy	Incretin-like weight loss, better tolerability, convenience, lower COGS with oral small molecule
2	NMRA-215 as add-on to a GLP1RA	Reduce incretin doses, increase response rate, enhance weight loss
3	NMRA-215 as weight maintenance treatment	Long-term cost effective and tolerable maintenance option

Potential to prove all hypotheses with higher-quality weight loss (e.g., preserving lean muscle mass)



Pre-Clinical Programs Each Have A Strong Biological Rationale

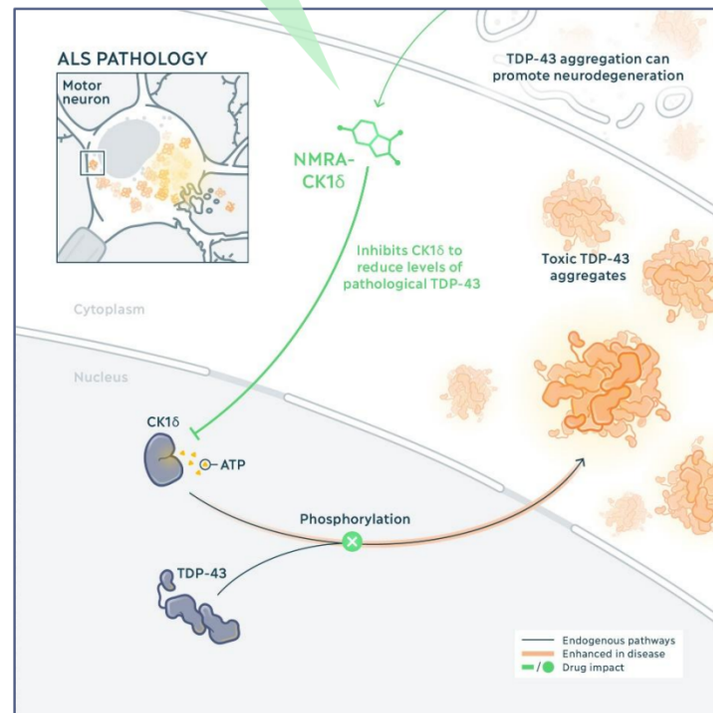
NMRA-CK1δ

Focused on inhibiting the protein casein kinase-1δ (CK1δ) to reduce levels of the pathological form of TDP-43 and slow disease progression in ALS

Potential Indications

ALS, Parkinson's disease

CK1δ phosphorylates TDP-43, a key driver of TDP-43-driven pathology in ALS



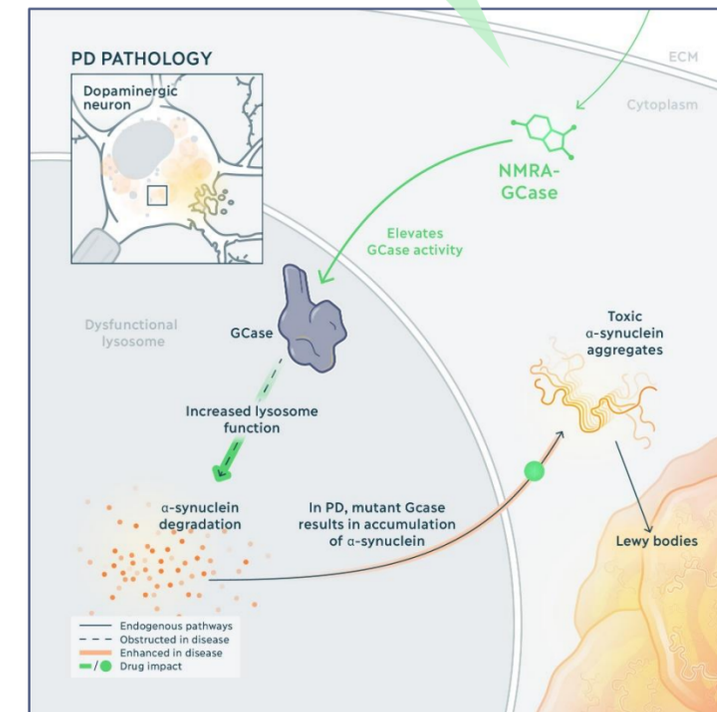
NMRA-GCCase

Focused on elevating activity of the GCCase enzyme, which is encoded by the GBA1 gene, and may help to degrade toxic a-synuclein aggregates

Potential Indications

Parkinson's disease

GCCase deficiencies lead to lysosomal dysfunction and the accumulation of alpha-synuclein, a hallmark of Parkinson's



Redefining Neuroscience Drug Development



**Industry leading
CNS pipeline with long-
dated IP into the 2040s**

**Multiple value-creating
clinical catalysts
on the horizon**



**Built at scale with strong
balance sheet; \$850M
raised since 2021**

**Cash runway into
2027 supporting
company growth**



**World-class team with
differentiated approach**

**Maximizing probability of
success with team and
proprietary approach**



Appendix



Led by Experienced Company Builders and Leading Neuroscience Drug Developers

Leadership



Paul L. Berns

Co-Founder, Chief Executive Officer & Chairman of Board of Directors



Joshua Pinto, Ph.D.

President



Bill Aurora, Pharm.D.

Chief Operating & Development Officer



Carol Suh

Chief Strategy Officer & Co-Founder



Jason Duncan

Chief Legal & Administrative Officer



Nick Brandon, Ph.D.

Chief Scientific Officer



Michael Milligan

Chief Financial Officer



Lori Houle

Chief Technical Operations & Quality Officer



Amy Sullivan

Chief Human Resources Officer



Pablo Gersberg

Chief Information Officer



Board of Directors

Paul L. Berns

Co-Founder, Chief Executive Officer, Chairman

Kristina Burow

Managing Director, ARCH Venture Partners

Matthew K. Fust

Biotechnology Advisor

Alaa Halawa

Executive Director, Mubadala Capital

Maykin Ho, Ph.D.

Retired Partner, Goldman Sachs

David Piacquad

Biotechnology Advisor



KOASTAL-1 Topline Study Summary Results

The KOASTAL-1 study enrolled 383 adult patients with Major Depressive Disorder (MDD)

Outcome	MADRS Total Score			SHAPS Total Score		
	Navacaprant 80 mg	Placebo	LSMD	Navacaprant 80 mg	Placebo	LSMD
ITT population CFB at Week 6 (Primary Endpoint)	-12.5 (n = 191)	-12.5 (n = 192)	0.0 (p = 0.993)	-5.8 (n = 191)	-5.5 (n = 192)	-0.3 (p = 0.648)
Female population CFB at Week 6	-14.0 (n = 105)	-11.4 (n = 106)	-2.7 (p = 0.072)	-7.2 (n = 105)	-4.9 (n = 106)	-2.3 (p = 0.015)
Male population CFB at Week 6	-10.6 (n = 86)	-13.8 (n = 86)	3.2 --	-4.3 (n = 86)	-6.3 (n = 86)	2.0 --

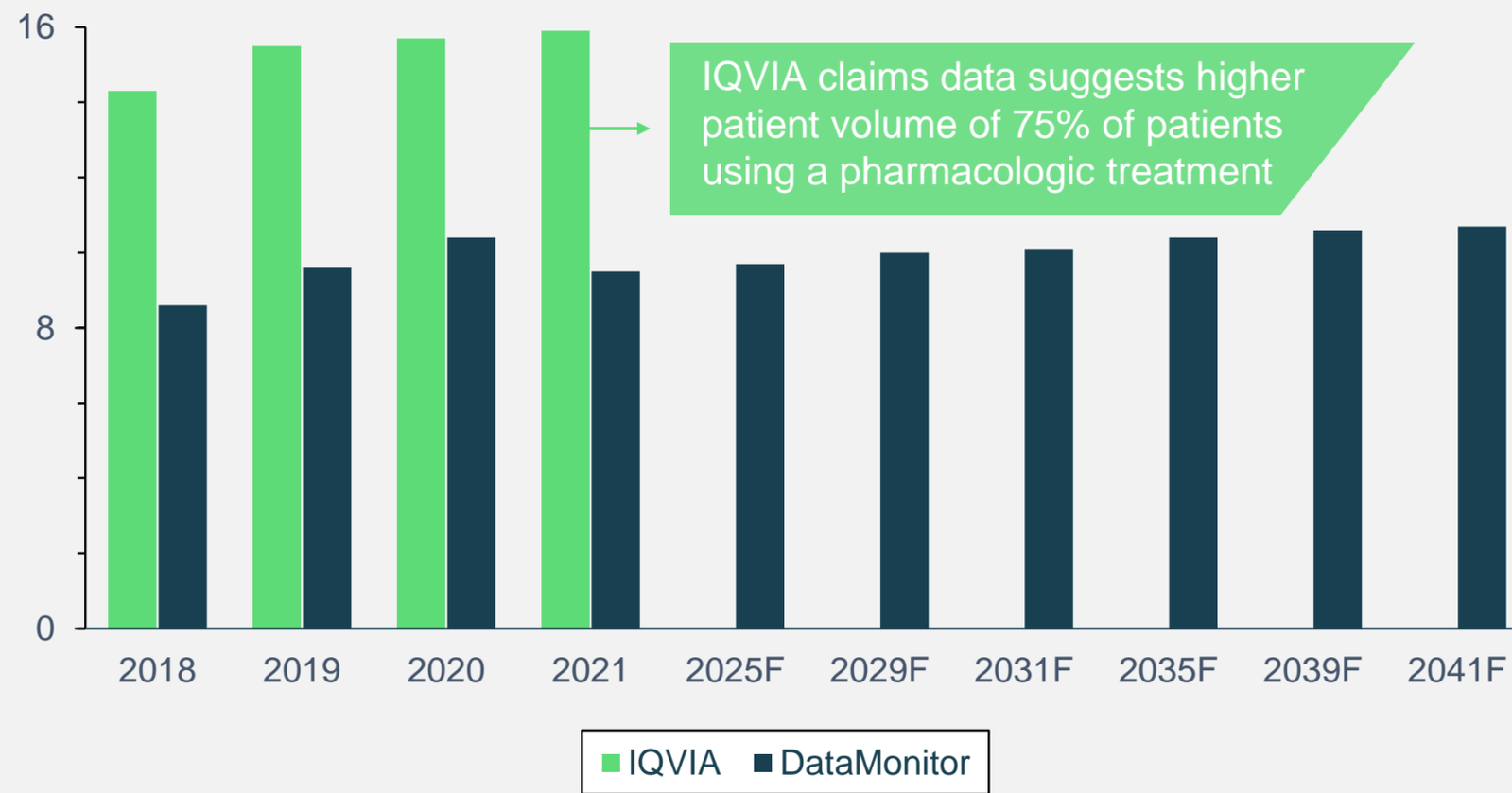
CFB = change from baseline; LSMD = difference in LS mean change from baseline between navacaprant and placebo groups generated from mixed-effects model for repeated measures. Subgroup analysis for male or female are pre-specified.



Navacaprant Would Enter Large MDD Market with a Highly Differentiated Profile

GROWTH IN ADDRESSABLE MDD MARKET EXPECTED IN-LINE WITH POPULATION GROWTH

U.S. MDD diagnosed, pharmacologically treated prevalent population (2018-41F) Millions of people



60-80% of MDD patients across lines of therapy are treated with a monotherapy agent¹

Monotherapy treatment rates across lines of therapy

Treatment Line	CCAIE	MDCD	MDCR	Optum
1 st	79.6%	82.1%	84.6%	81.7%
2 nd	67.3%	67.8%	69.3%	66.1%
3 rd	63.9%	64.9%	67.2%	62.1%
4 th	61.4%	61.4%	68.1%	60.0%

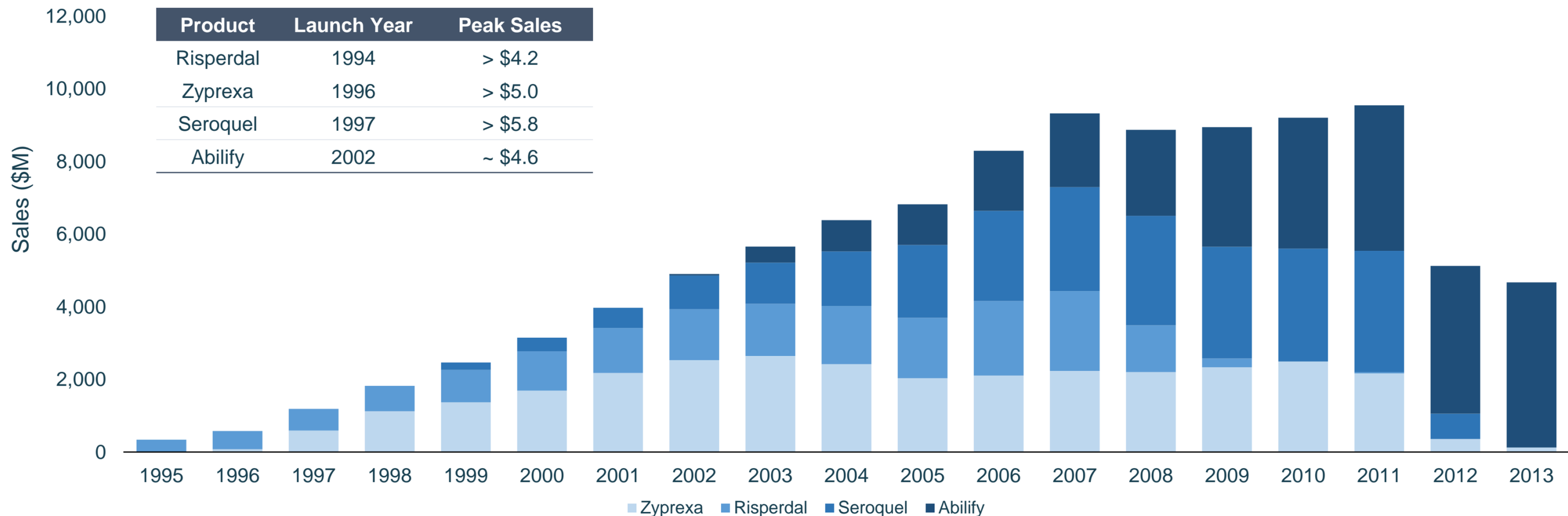
¹Kern et al. Treatment patterns and sequences of pharmacotherapy for patients diagnosed with depression in the United States: 2014 through 2019. BMC Psychiatry. (2020) 20:4. U.S. Census Population Projections; DRG; Datamonitor; National Survey of Drug Use and Health 2018, 2019, 2020, 2021; Torre et al. (2021); L.E.K. research and analysis CCAIE = IBM MarketScan Commercial Database; MDCD = IBM Market Scan Multi-State Database; MDCR = IBM MarketScan Medicare Supplemental Database



Schizophrenia Market Supports Multiple Treatment Options

Historically the schizophrenia market has supported multiple branded products with similar MOAs, with new entrants driving higher overall market sales volume

Sales of Branded 5-HT2 to D2 Receptor Antagonists (1995 – 2013)



Sources: EvaluatePharma, L.E.K. interviews, research, and analysis; GK associates "The order of entry effect in prescription (Rx) and over the counter (OTC) pharmaceutical drugs", International Journal of Pharmaceutical and Healthcare, Marketing Vol. 2 No. 1, 2008 pp. 35-46. MOA = Mechanism of Action.

Opportunity to Build a Leading Neuropsych Product Franchise

Potential for **broad indication expansion** in poorly served disorders with a **novel mechanism**

