



Transforming the Possible in Neuroscience

4Q 2020 Financial Results & Pipeline Update

March 2021



Forward-Looking Statements

This presentation contains forward-looking statements that are based on management's beliefs and assumptions and on information currently available to management. In some cases, you can identify forward-looking statements by the following words: "may," "will," "could," "would," "should," "expect," "intend," "plan," "anticipate," "believe," "estimate," "predict," "project," "potential," "continue," "ongoing" or the negative of these terms or other comparable terminology, although not all forward-looking statements contain these words. These statements involve risks, uncertainties and other factors that may cause actual results, levels of activity, performance or achievements to be materially different from the information expressed or implied by these forward-looking statements. Although we believe that we have a reasonable basis for each forward-looking statement contained in this presentation, we caution you that these statements are based on a combination of facts and factors currently known by us and our projections of the future, about which we cannot be certain.

Forward-looking statements in this presentation include, but are not limited to: statements about the potential attributes and benefits of our product candidates, including with respect to receptor subtype selectivity, activity, side effect and tolerability profile and relevant indications; the format and timing of our product development activities and clinical trials, including the design of clinical trials and the timing of initiation, completion and data readouts for clinical trials; the timing and outcome of IND submissions and other regulatory interactions; the ability to compete with other companies currently marketing or engaged in the development of treatments for relevant indications; the size and growth potential of the markets for product candidates and ability to serve those markets; the rate and degree of market acceptance of product candidates, if approved; the potential effects of the business combination; cooperative grant funding from NIDA; and the sufficiency of our financial resources.

We cannot assure you that the forward-looking statements in this presentation will prove to be accurate. Furthermore, if the forward-looking statements prove to be inaccurate, the inaccuracy may be material. Actual performance and results may differ materially from those projected or suggested in the forward-looking statements due to various risks and uncertainties, including, among others: that clinical trial results may not be favorable; uncertainties inherent in the product development process (including with respect to the timing of results and whether such results will be predictive of future results); the impact of COVID-19 on the timing, progress and results of ongoing or planned clinical trials; other impacts of COVID-19, including operational disruptions or delays or to our ability to raise additional capital; whether and when, if at all, our product candidates will receive approval from the FDA or other regulatory authorities, and for which, if any, indications; competition from other biotechnology companies; uncertainties regarding intellectual property protection; and other risks identified in our SEC filings, including those under the heading "Risk Factors" in our prospectus filed with the SEC on December 4, 2020 and our subsequent SEC filings.

In light of the significant uncertainties in these forward-looking statements, you should not regard these statements as a representation or warranty by us or any other person that we will achieve our objectives and plans in any specified time frame, or at all. The forward-looking statements in this presentation represent our views as of the date of this presentation. We anticipate that subsequent events and developments will cause our views to change. However, while we may elect to update these forward-looking statements at some point in the future, we have no current intention of doing so except to the extent required by applicable law. You should, therefore, not rely on these forward-looking statements as representing our views as of any date subsequent to the date of this presentation.

Agenda

Introduction



Matthew Calistri
Vice President, Investor Relations

Overview



Tony Coles, M.D.
Chairperson & Chief Executive Officer

Lead Program Updates



Raymond Sanchez, M.D.
Chief Medical Officer

Early Pipeline Updates



John Renger, Ph.D.
Chief Scientific Officer

Financial Performance & Outlook



Kathy Yi
Chief Financial Officer

Q&A

All

Company Highlights



Pursuing a targeted approach to neuroscience that combines a differentiated understanding of neurocircuitry with advanced chemistry to develop novel therapies for CNS diseases



Broad portfolio of 11 assets targeting large markets with significant unmet need, including schizophrenia, epilepsy and Parkinson's Disease



Progressing towards multiple near and medium-term catalysts, with up to 8 data readouts and multiple INDs expected by the end of 2023



Leveraging a seasoned management team with extensive expertise in neuroscience and a strong track record of over 20 prior drug approvals and commercialization



2020 Highlights and 2021+ Outlook

Corporate Updates

- Completed go-public transaction - **\$440 million net proceeds**
- Opening **new headquarters** at Cambridge Crossing

Significant Pipeline Progress

- Initiated dosing in all **6 ongoing clinical trials** and 2 open-label extensions*
- **Submitted IND** for CVL-871 in dementia-related apathy
- Expect to receive **NIDA cooperative grant funding** for CVL-936 (D3 Antagonist)

Upcoming Data Readouts

- **CVL-231:** Phase 1b in Schizophrenia (**mid-year 2021**)
- **Darigabat:** Phase 1 in Acute Anxiety (**2H 2021**) & Phase 2 in Focal Epilepsy (**2H 2022**)
- **Tavapadon:** Phase 3 in Late PD (**1H 2023**) & Early PD (**2H 2023**)
- **CVL-871:** Phase 2a in Dementia-Related Apathy (**2H 2022**)

Financial Outlook

- Increased R&D spending to **ramp-up clinical trials** & invest in early pipeline
- Cash balance of **\$384 million** as of 12/31/2020
- Cash runway projected into **2023**



\$350 million capital commitment



Portfolio built on:
10+ years of research
\$1 billion+ R&D investment

Founded 2018

Perceptive Advisors & Others



Pre-\$: \$780M
Post-\$: ~\$1.3B

SPAC Transaction Completed 2020

11 Neuroscience assets

Novel targets; small molecules

Experienced management team

Development driven by data

Led by a Seasoned Life Sciences Management Team



Tony Coles, M.D.
*Chairperson &
Chief Executive Officer*



Kathy Yi
Chief Financial Officer



Raymond Sanchez, M.D.
Chief Medical Officer



John Renger, Ph.D.
Chief Scientific Officer



Kenneth DiPietro
*Chief Human
Resources Officer*



Kathleen Tregoning
*Chief Corporate
Affairs Officer*



Strong Track Record of Approvals



Cerevel's Targeted Approach to CNS Disease

Leveraging Expertise in Neurocircuitry

Pipeline Uniquely Based on

Differentiated Understanding of Neurocircuitry

Targeted Receptor Subtype Selectivity

Receptor Binding/Modulation

Optimized Receptor Pharmacology

Highly Selective Small Molecules in Clinical Studies
Created using Pfizer world-class chemistry

Robust Data Packages

Deep Pipeline: Multiple Value Inflections Near & Long-Term

11 Assets

5
Clinical Assets

Tavapadon

Darigabat

CVL-231

CVL-871

CVL-936

4
Pre-Clinical Assets

LRRK2*

M4 Agonist*

PDE4B*

KORA (CVL-354)

2
Undisclosed Assets

Patient Populations

Early Parkinson's

Late Parkinson's

Epilepsy

Anxiety

Schizophrenia

Dementia-Related
Apathy

Substance Use Disorder

8 Trials

3 Phase 3 Trials

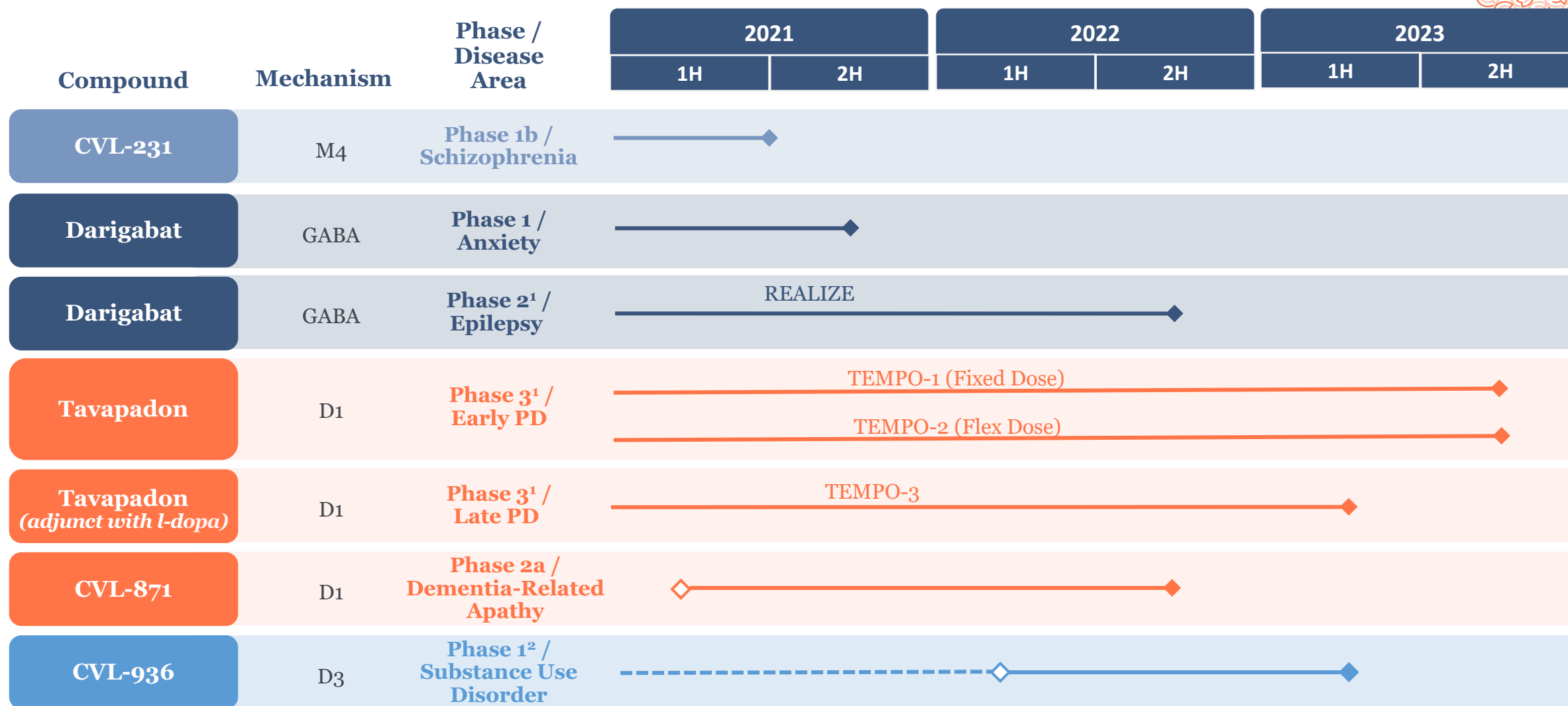
2 Phase 2 Trials

1 Phase 1b Trial

2 Phase 1 Trials

+2 Open Label Extensions

Multiple Milestones Expected Over Next Three Years

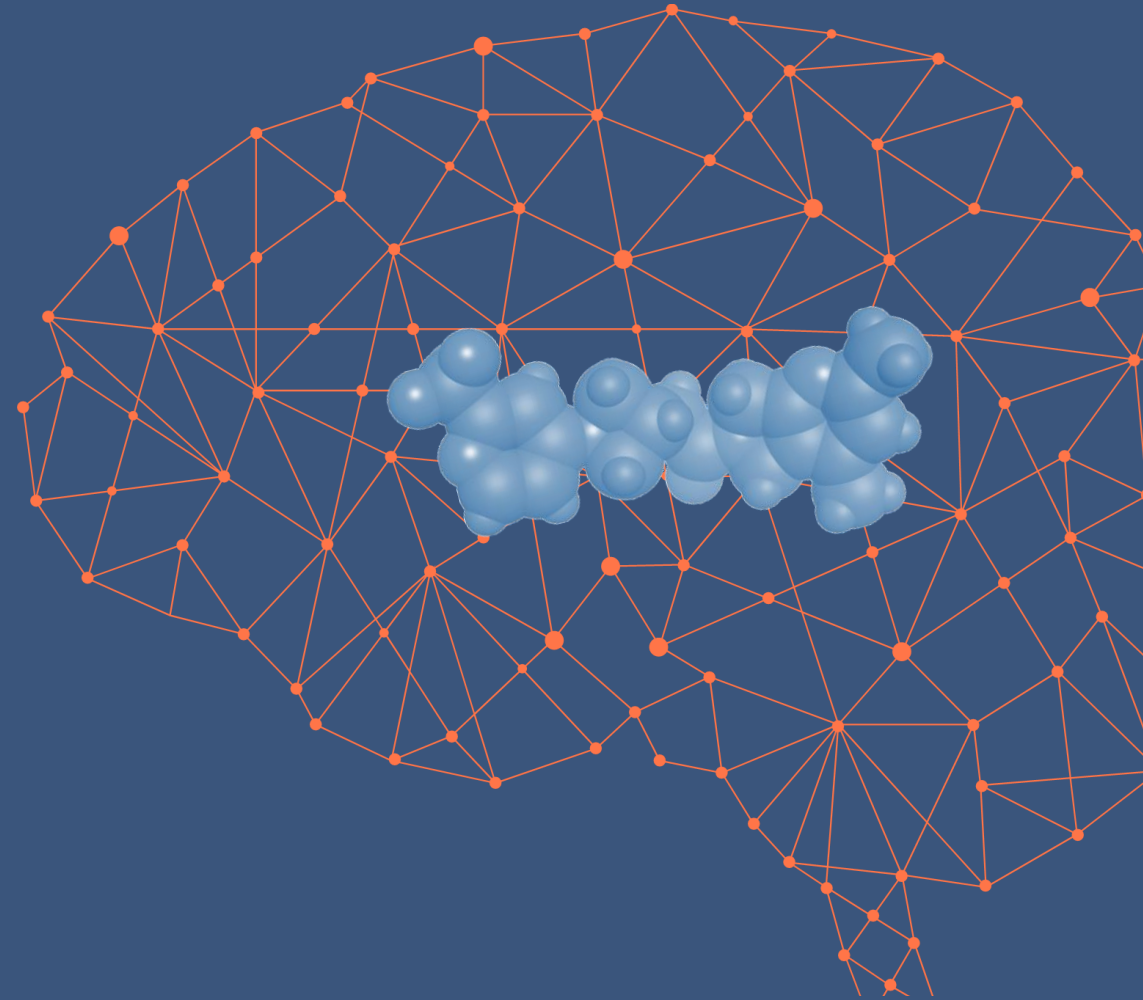


◇ Estimated Trial Initiation

◆ Estimated Topline Data

■ M4 PAM (CVL-231) in Schizophrenia

Selectively targeting the M4 muscarinic receptor with the goal of treating psychosis-related symptoms with improved side effect profile



Cerevel's M4 PAM is a Potential Next Generation Antipsychotic

M4 PAM (CVL-231)

Potential New Standard of Care

**First-in-Class Therapy
with Novel MOA**

M4 Selective

Targeted Muscarinic Activity

Improved Tolerability

Opportunity for Innovation in Schizophrenia

Current Standard of Care Uses Same
Mechanism of Action (MOA) as Therapies from the 1950s

Large
Market

~21M
Patients
Worldwide

>\$9B
Revenues
in 2018

~3.5%
Growth
per year

Significant
Need for New
Treatment
Option

Side Effect and
Tolerability Issues

High Discontinuation
74%
Within 18 months

Limited
Compliance

60%

Progression and
worsening of disease

High
Relapse Rates

77%

at 1 year

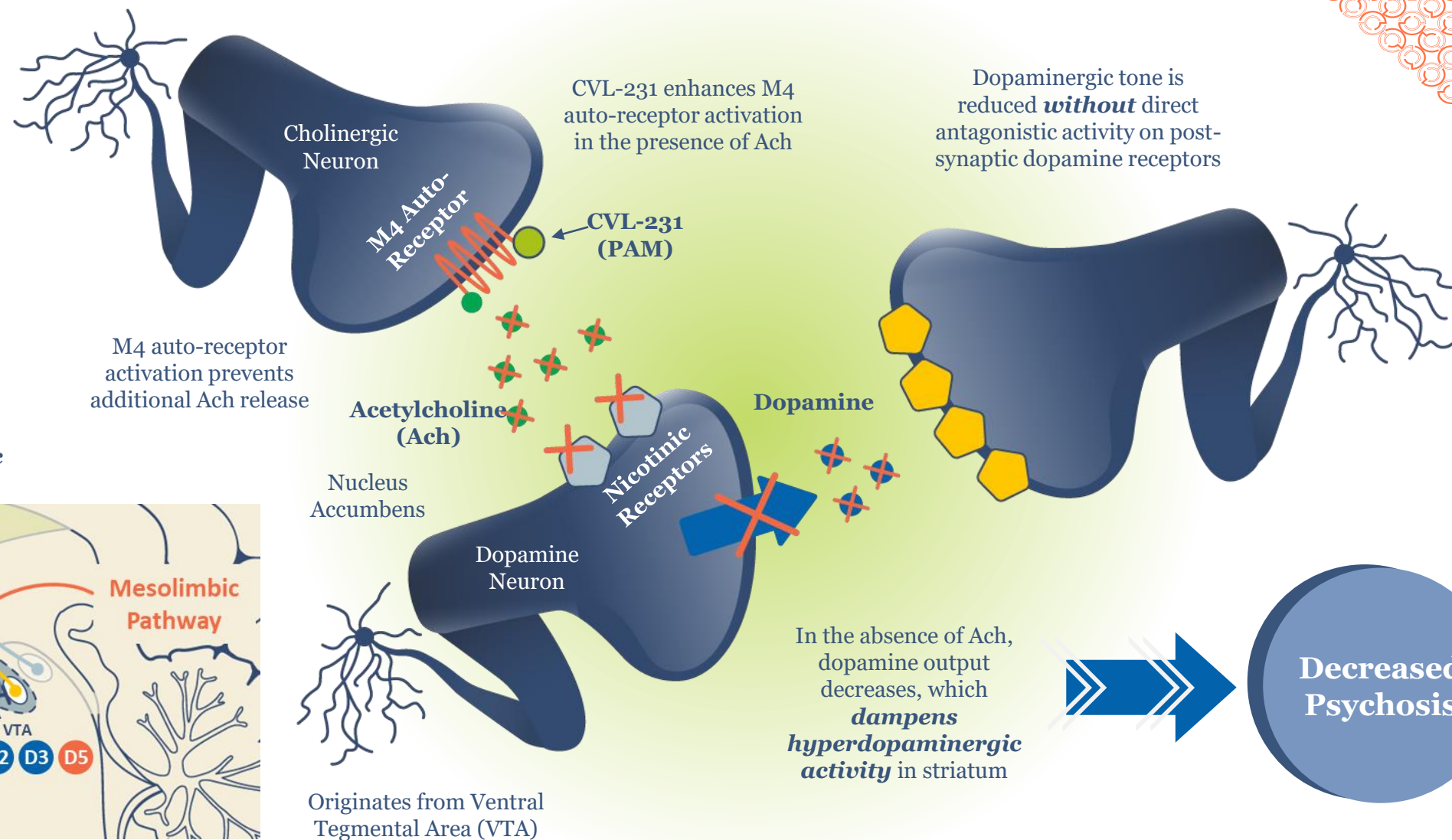
90%

at 2 years

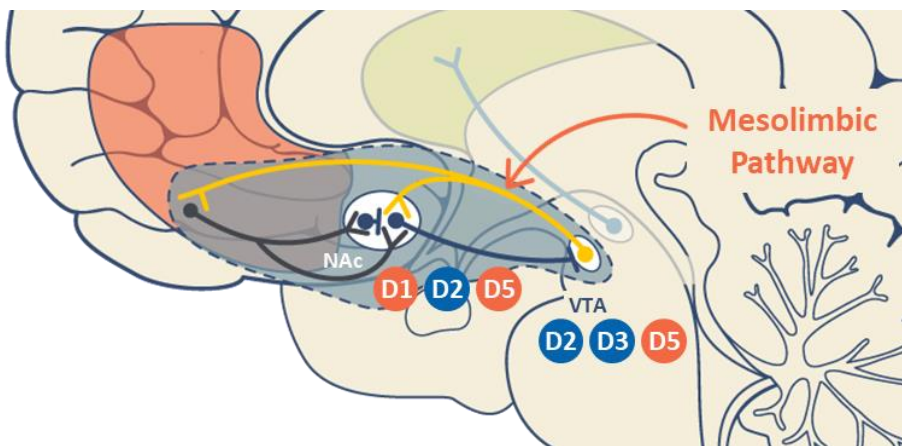
Debilitating side effects of atypicals often lead to discontinuation and relapse,
driving a vicious cycle of disease progression



M4 Receptor Activation Reduces Dopamine in the Striatum



Schizophrenia symptoms driven by **overactivity of the dopaminergic mesolimbic pathway**



Cerevel's Selective M4 Modulation: A Compelling and Differentiated Approach to Drive Antipsychosis

M4 Selectively Impacts Brain Functions

Other Muscarinic Receptors	Potential Effect	M4 Muscarinic Receptor
-	Antipsychosis	✓✓
✓✓	Cognition	✓
✓✓	GI Side Effects	-
✓	Cardiovascular	✓

Receptor Subtype Selectivity Offers Potential Improvement

Xanomeline (M1/M4) data from Schizophrenia and Alzheimer's patients show targeting muscarinic receptor impacts brain function
But development limited by GI and CV side effects

Karuna's KarXT creatively addresses this by adding trospium to Xanomeline to offset side effects
Non-selective approach

M4 Knock-out mouse data suggests M4 receptors drive the antipsychotic activity of Xanomeline
M1 receptors believed to contribute to worrisome side effects

CVL-231:
Selective Potentially Once-daily M4 PAM

>600X
more selective for
M4 over M1, 3 and 5

~360X
more selective
than for M2



Source: 1. Shekhar et al. Am J Psychiatry, Vol 165, Aug 2008. N=20. 2 active and 3 placebo-arm patients discontinued the study, none due to adverse events.
2. Bodick et al. Arch Neurol, Vol 54, Apr 1997. N = 343. 52% of patients in high-dose arm discontinued treatment due to adverse events.

M4 PAM Ongoing and Planned Studies

Study 001 – Phase 1b

Part A: Safety Assessment

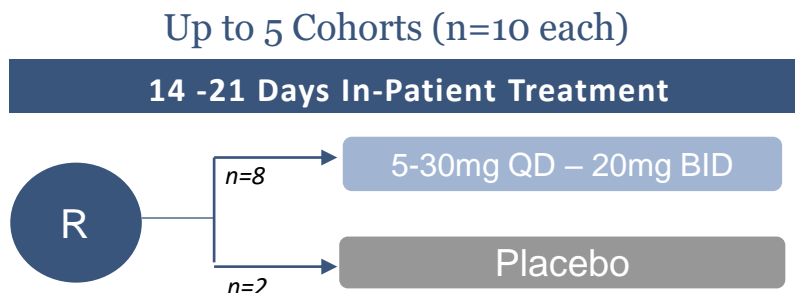
Multiple Ascending Dose

Primary Objective

- Safety & tolerability

Secondary Objective

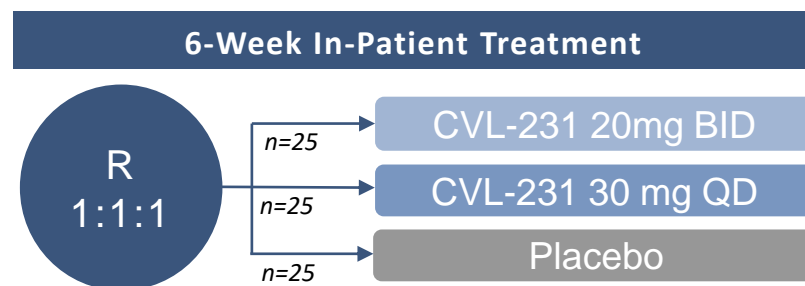
- PK



Part B: Pharmacodynamics

Exploratory PD Assessment

- Positive and Negative Syndrome Scale (PANSS)*
- Clinical Global Impression – Severity Scale (CGI-S)*
- Brief Assessment of Cognition in Schizophrenia (BACS) symbol coding test*



PET Studies

Study 002 – Phase 1b

Single Dose (n=9)

Designed to inform PK vs. target receptor occupancy

Study Objectives

Primary
M4 CNS receptor occupancy vs. peripheral drug exposure

Secondary
Safety and tolerability

Study 003 – Phase 1b

Single Dose (n=9)

Designed to inform receptor occupancy vs. target pharmacology

Study Objectives

Primary
Modulation of striatal levels of dopamine with CVL-231

Secondary
Safety and tolerability



Data for Phase 1b Trial Expected Mid-Year 2021

Dementia-Related Psychosis (DRP): Potential Opportunity for CVL-231 Beyond Schizophrenia

DRP Overview and Unmet Needs¹⁻⁷

- Psychosis incidence ranges from 10-75% of Alzheimer's patients and varies by stage of disease
 - Upwards of 1M moderate to severe Alzheimer's patients in the G7 experience symptoms of psychosis
- Co-morbidities including agitation, aggression and depression
- Often leads to long-term care / nursing home admissions

Standard of Care

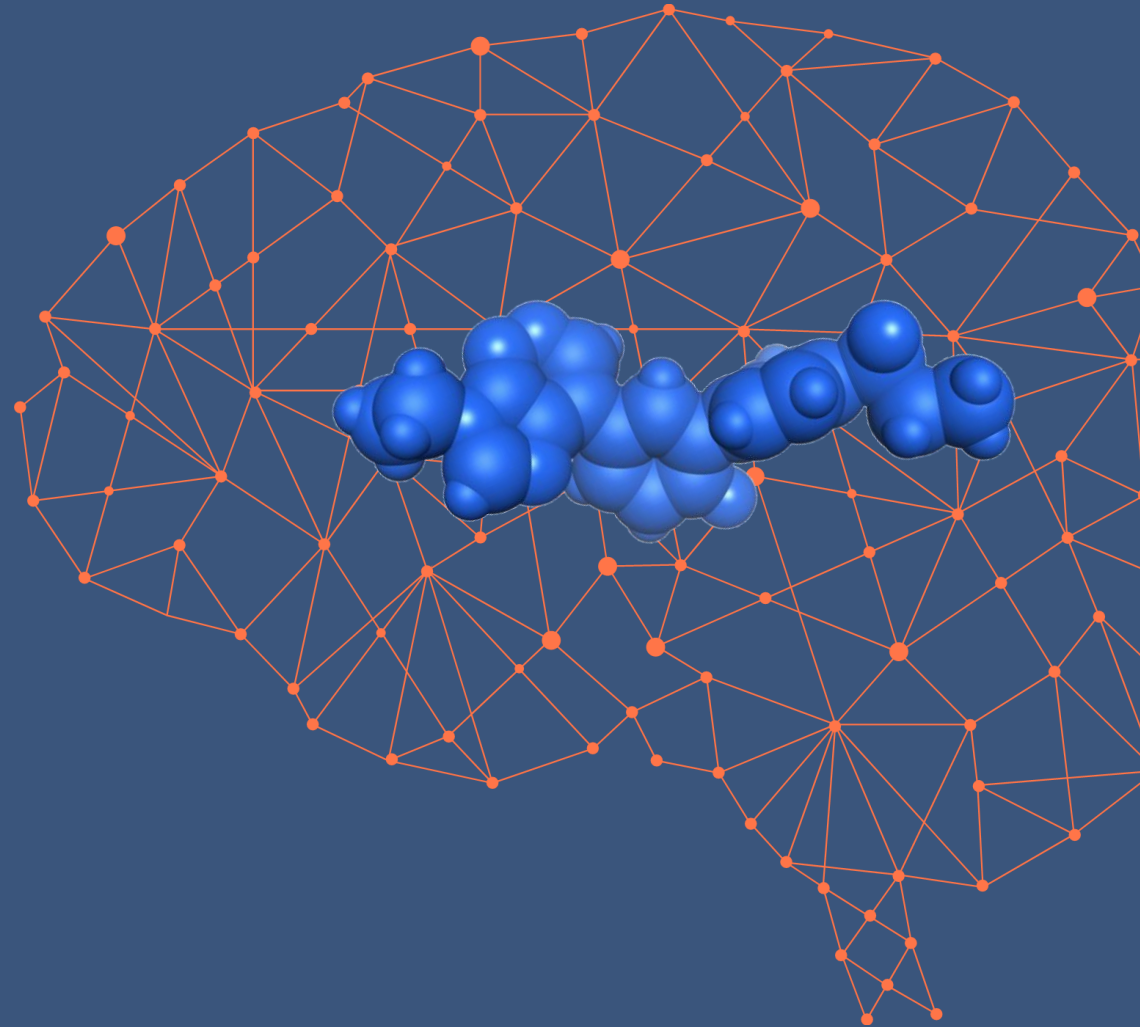
- None established
- Off-label use of atypical antipsychotics: tolerability issues heightened in this population; contribute to cognitive decline

Next Steps for CVL-231

- CVL-231 side effects / tolerability observed to date are appropriate for further clinical evaluation in elderly patients
- Upcoming clinical pharmacology study in the elderly

■ Darigabat (GABA_A PAM)

Selectively targeting the α -2/3/5 subunits of the GABA_A receptor with the goal of enhancing anti-convulsant and anxiolytic effects without dose-limiting sedation



Darigabat has Potential for Benzo-like Activity, Improved Side Effects and Chronic Dosing

Darigabat

Potential to become first-line and adjunct therapy

Targeted GABA α 2/3/5 Receptor Selectivity

Benzo-like Activity

Improved Tolerability

Potential for Reduced Abuse Liability

Opportunity for New Treatment Option in Epilepsy

HCPs and patients are dissatisfied due to insufficient activity, side effects and poor tolerability

Large Market

▶ ~65M Patients Worldwide

>\$6B G7 Revenues in 2018

~6% per year Branded AED¹ Market Growth through 2025

Benzos are highly efficacious, but...

▶ Poor Tolerability

Desensitization & Loss of Efficacy

Potential for Abuse

Withdrawal



Potential as chronic therapy with improved side effect profile and tolerability may expand use vs. traditional benzodiazepines

Selective GABA_A Receptor PAM: Potential for Lower Seizure Rates, Reduced Side Effects

GABA α -2/3/5 Can Differentially Address Symptoms

GABA subtype predicted effects:	α 1	Darigabat		
		α 2	α 3	α 5
Anti-convulsant	✓✓	✓✓		
Anxiolysis		✓✓	✓✓	
Analgesia		✓✓	✓	✓✓
Muscle Relaxation		✓✓	✓✓	
Sedation	✓✓			
Cognitive Impairment	✓✓	?	?	✓
Addiction	✓✓	✓		

Benzodiazepine
side effects

Role for Targeted GABA α 2/3/5 Receptor Selectivity

- >10 benzodiazepines (BZDs - broad-spectrum GABA modulators) widely prescribed for anxiety, seizures and other indications
- Significant need for better-tolerated, less sedating and less addictive GABA modulators
- BZD Rx's increased >95% from 2003-2015



To our knowledge, darigabat is the only GABA α -2/3/5 selective PAM in clinical trials for epilepsy

Darigabat Data Showed a Favorable Side Effect Profile Relative to Benzodiazepines

Multiple doses of darigabat

Phase 1 MAD Study – only mild AEs and limited somnolence up to 42.5 mg BID

Able to achieve >80% receptor occupancy without significant somnolence observed

Benzodiazepines achieve 10% to 15% receptor occupancy with significant somnolence observed

No evidence of withdrawal effects

Phase 1 MAD Study (Protocol: B7431011)

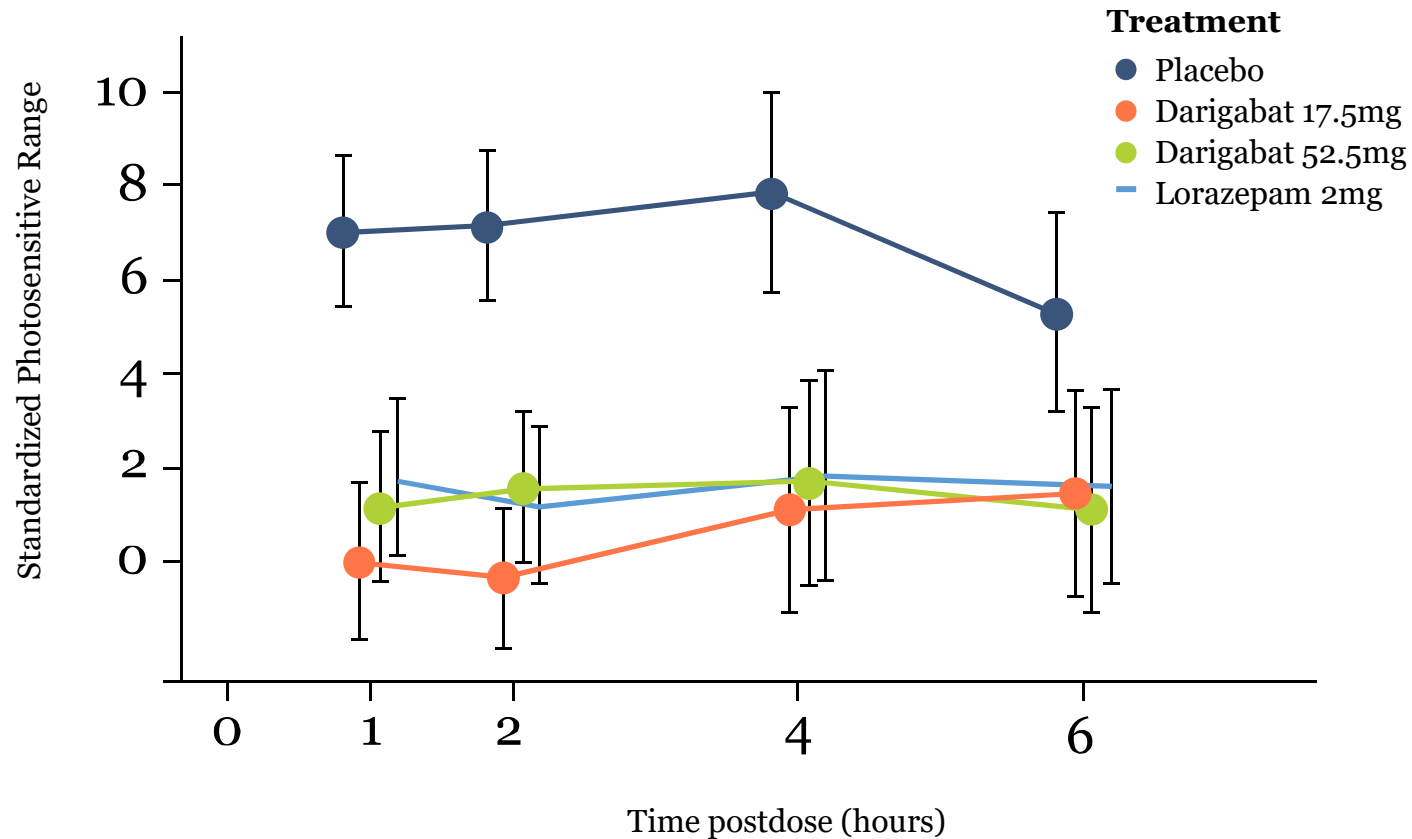
	Reaction	Week 1 (Titration)	Week 2 (Maintenance)	Week 3 (Maintenance)	Follow-up
Placebo	No Reaction	4 / 4	4 / 4	3 / 4	4 / 4
	Dizziness	-	-	1 / 4	-
	Somnolence	-	-	-	-
25 mg BID (~80% RO ⁽¹⁾)	No Reaction	5 / 8	7 / 8	8 / 8	8 / 8
	Dizziness	2 / 8	1 / 8	-	-
	Somnolence	3 / 8	-	-	-
42.5 mg BID (>80% RO ⁽¹⁾)	No Reaction	4 / 7	6 / 7	6 / 7	6 / 7
	Dizziness	3 / 7	1 / 7	1 / 7	1 / 7
	Somnolence	-	-	-	-



No somnolence observed following titration through doses of 42.5 mg BID

Darigabat Phase 2 Data Showed Benzo-like Anticonvulsant Activity in Photosensitive Epilepsy⁽¹⁾

Darigabat in Single-Dose Photosensitive Epilepsy Study



Darigabat Results

Anticonvulsant activity comparable to lorazepam

Improved sedation and AE profile compared to benzos

Complete suppression in 6 of 7 subjects

Majority of AEDs developed for epilepsy that showed positive published photoepilepsy results were approved⁽²⁾

Darigabat REALIZE Trial: Data Expected 2H 2022

REALIZE Phase 2 Trial In Focal Epilepsy

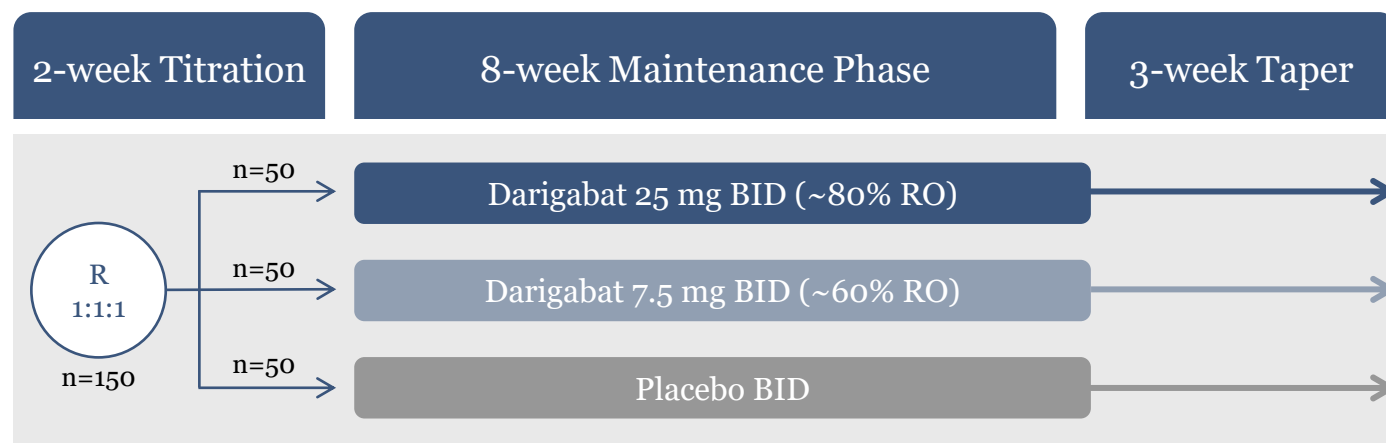
Targeting ~60 sites in 3 countries

Inclusion criteria

- Adults (18-75) with drug-resistant focal onset seizures
- History of 4+ seizures per month for at least 3 months
- 1-3 stable background AEDs allowed

Primary endpoint

- Reduction in focal onset seizure frequency



Patients able to join 57-week open-label extension trial (REALIZE OLE) after completion of 8-week maintenance phase



Focal epilepsy trial intended to establish proof of concept and side effect profile to support development in broader epilepsy indications

Darigabat: Phase 1 Program in Acute Anxiety

The Hypercapnia (CO₂ Inhalation) Model

- CO₂ inhalation challenge is translational model providing proof-of-principle for anxiolytic activity in early clinical development
- Well-established in both healthy volunteers and in patients with panic disorder
 - Hypercapnia results in increased fear and panic, as measured by Visual Analogue Scales (VAS) and the Panic Symptom List (PSL)¹
- The proposed mechanism of the anxiety induced by hypercapnia model is decreased GABA and increased noradrenaline²
- The model is sensitive to drugs used to treat anxiety disorders (including benzodiazepines & SSRIs) and emerging new treatments with novel mechanisms

CO₂ inhalation induces fear and panic symptoms in healthy volunteers and panic disorder patients

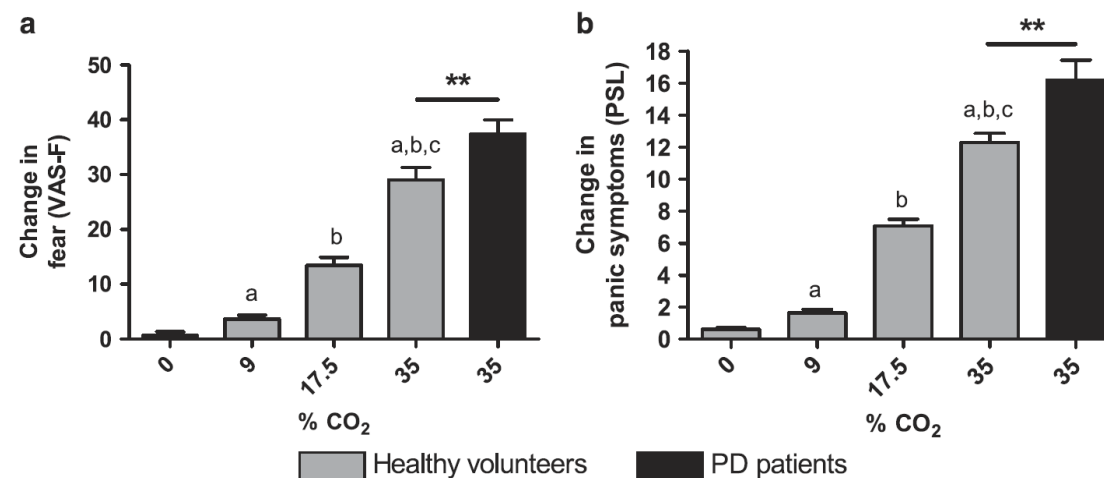
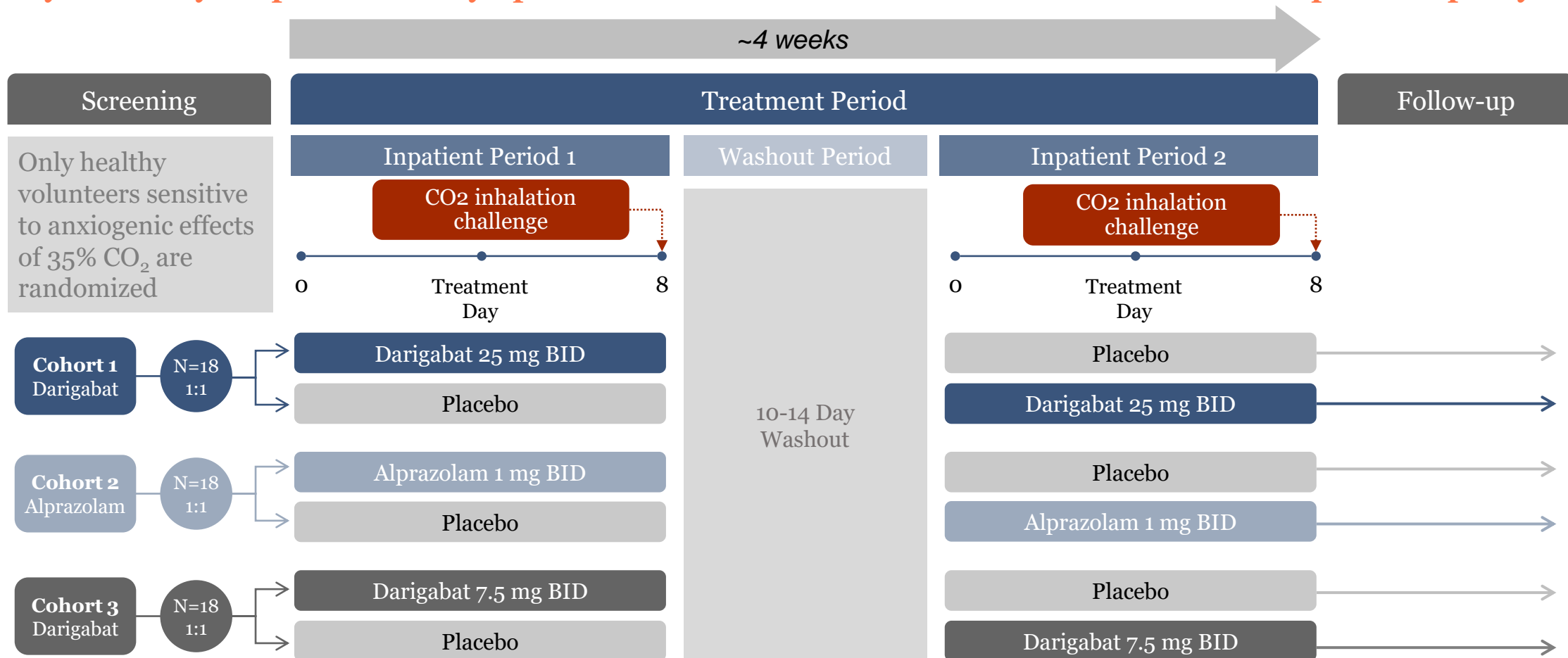


Figure 2. Effect of CO₂ on self-reported fear and panic symptoms in healthy volunteers and PD patients. In healthy volunteers (gray), both fear (a) and panic symptoms (b) increased dose-dependently. Inhaling 35% CO₂ triggered a more robust response in patients (black) when compared with healthy volunteers. Data represent mean+s.e.m. (a) Compared with 0% CO₂, $P < 0.001$; (b) compared with 9% CO₂, $P < 0.001$; (c) compared with 17.5% CO₂, $P < 0.001$; ** $P < 0.01$. PD, panic disorder; PSL, Panic Symptom List; VAS-F, Visual Analog Scale for fear.

Darigabat Phase 1 in Acute Anxiety: Data Expected 2H 2021

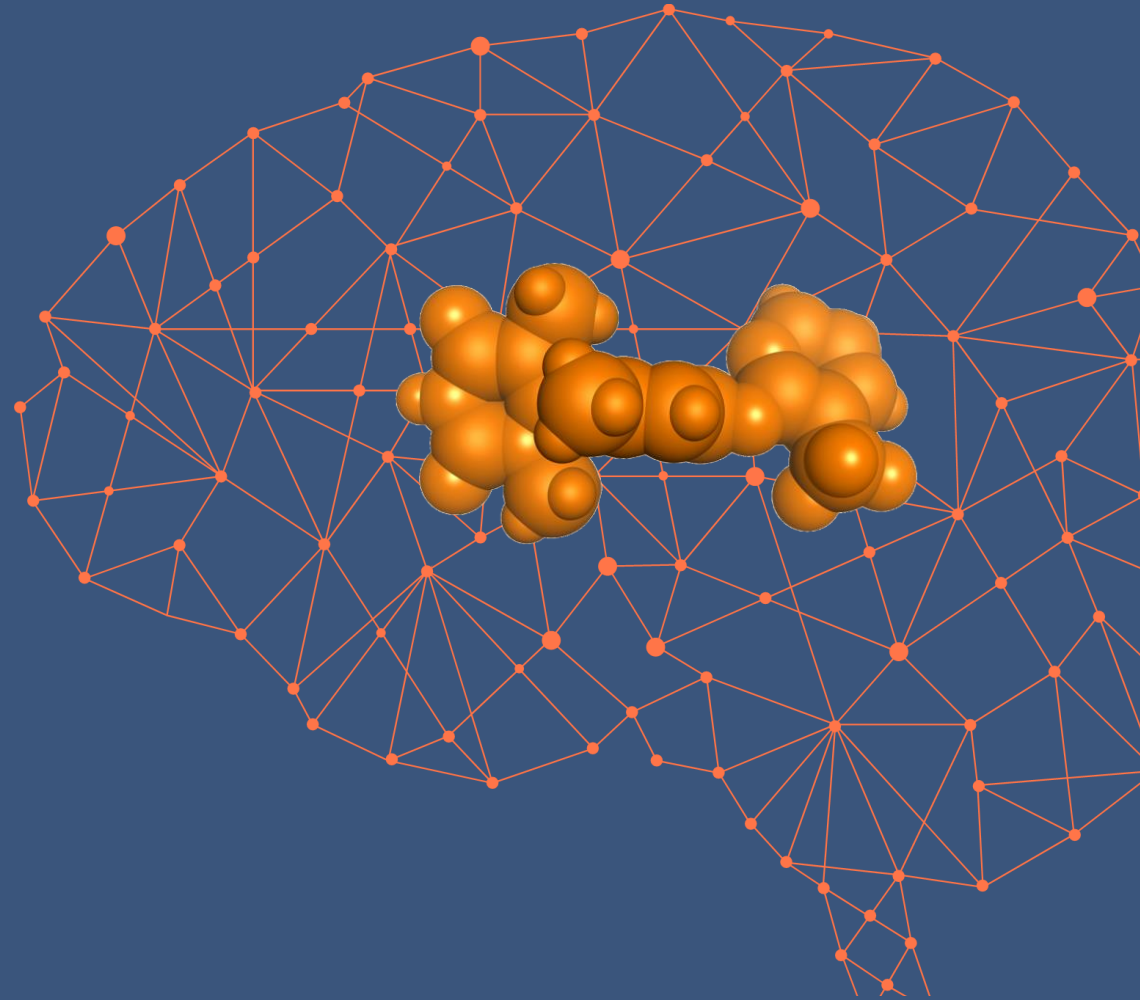
Randomized, double-blind, placebo- and active-controlled crossover design with multiple doses over 8 days. Primary endpoint: Panic symptoms list¹. Doses selected to achieve ~60 and 80% receptor occupancy



1. The Panic Symptom List (PSL) includes 13 symptoms scored across a range of 0 (absent) to 4 (very intense) that is used to assess panic anxiety. Liebold et al. Trans Psychiatry. 2016.; Bailey et al. J Psychopharm. 2011.; Malizia et al. Arch Gen Psychiatry. 1998.; Salvatore et al. Translational Psychiatry 2020.

D Tavapadon in Parkinson's Disease

Partial agonist selectively targeting the dopamine D1 receptor with the goal of enhancing motor control while minimizing side effects





Tavapadon has Potential to be a Differentiated Treatment for Parkinson's

Designed to be a novel backbone therapy for patients from diagnosis to the end of treatment:

Only* D1/D5 selective molecule

Avoid D2/D3 Side Effects: *Sudden daytime somnolence, hallucinations, acute orthostasis and impulse control disorders*

First* partial agonist for Parkinson's

Avoid Dyskinesias: *Driven by receptor overexcitation*

Predictable 24-hour activity

Sustained Effect: *Once daily, oral dosing*

Selective direct motor pathway activation

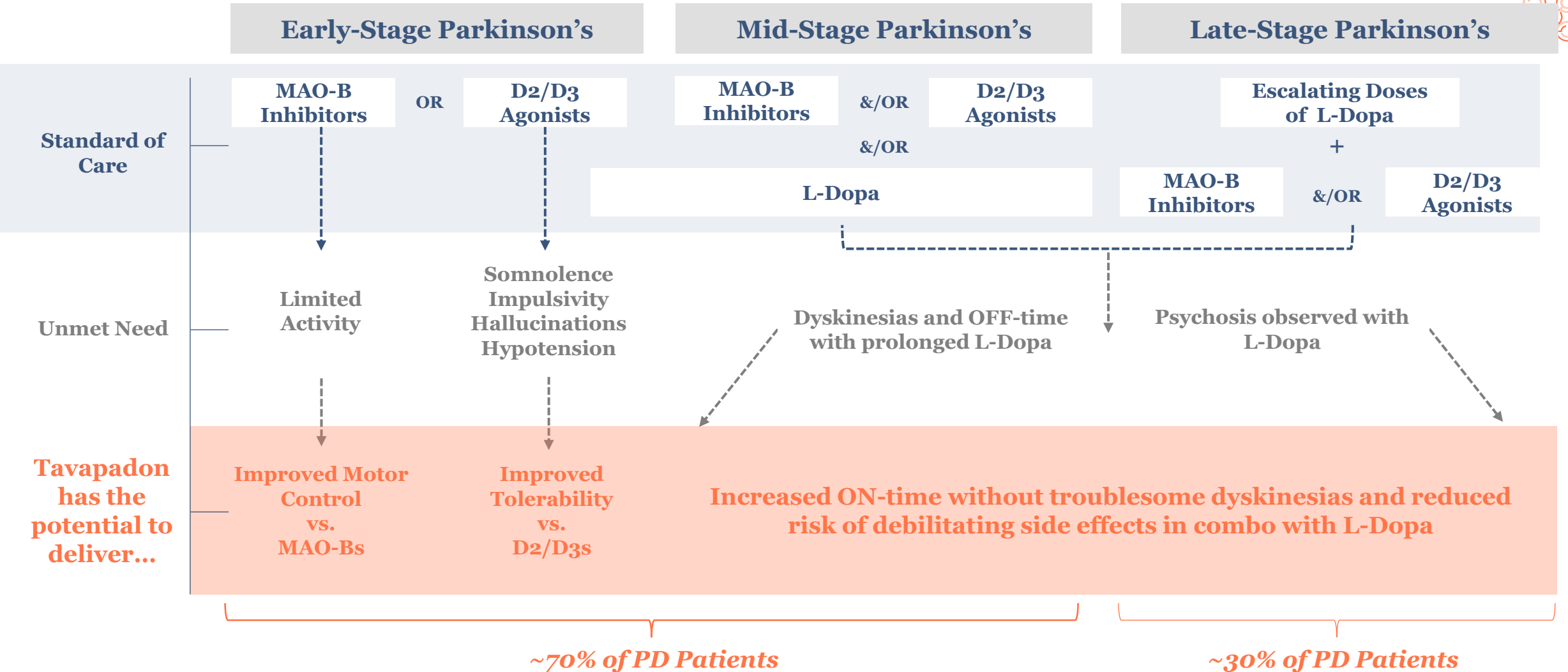
Superior motor control over D2/D3s full agonists

- Feedback received from FDA on our registrational program (2019)
- To our knowledge, nothing else in the symptomatic pipeline positioned to provide broad therapeutic benefit and differentiation



First-in-class potential designed to offer stable motor control and favorable side effect profile with broad monotherapy and adjunct therapy benefit

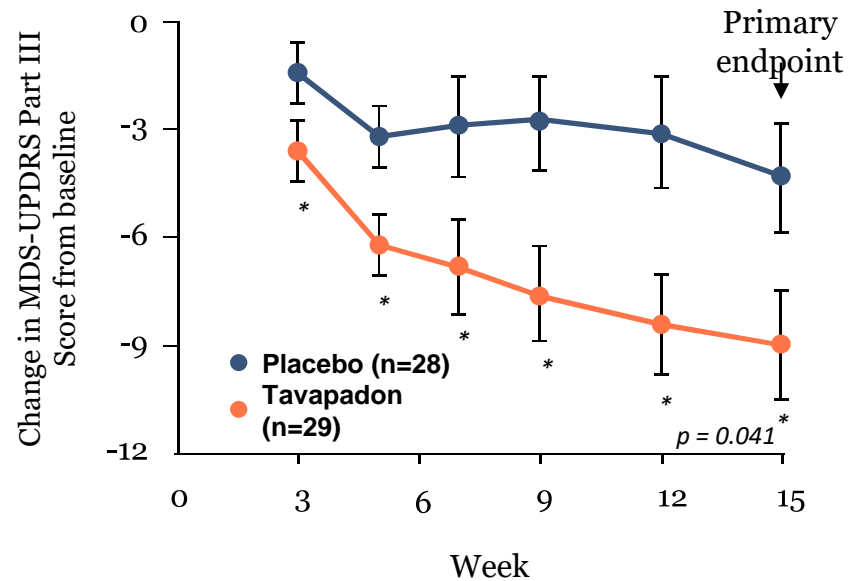
Tavapadon Designed to Address Unmet Needs Across All Stages of Parkinson's: Early and Late



Selective Direct Motor Pathway Activation Designed to Provide Differentiated Treatment Option in Early Parkinson's

Potential for motor control as good or better than D2/D3s with once-daily dosing and improved side effect profile

Phase 2 Data: Tavapadon in Early PD¹ (Primary Endpoint: MDS-UPDRS III Motor Score)



In Phase 2, tavapadon demonstrated 4.8 point MDS-UPDRS III difference vs. placebo at week 15 (p=0.04, MMRM)

Note: Average dose of tavapadon = 9 mg; 11 patients on 15 mg top dose at week 15. Baseline Part III scores of 24.3 (tavapadon) and 25.8 (placebo).

Additional Tavapadon Phase 2 Data¹

- When adjusted to exclude patients with baseline MDS-UPDRS II of 0 or 1, **showed improvement of ~2 points over placebo on MDS-UPDRS Part II²**
- Most common AEs included headache and nausea (can be mitigated with titration)
- Tavapadon's incidence of known D2/D3 side effects:
 - Somnolence: 14%
 - Nausea: 31%
 - Hallucinations: 0%³
 - Hypotension-Related Events: 7%
 - Dizziness: 7%

Tavapadon demonstrated 5.8 point improvement over placebo at week 15 on MDS-UPDRS Part II + III (p = 0.02, MMRM)

Tavapadon TEMPO-1 & -2 in Early PD: Data Expected 2H 2023



Registration-directed program includes three global Phase 3 trials optimally designed to maximize treatment effect

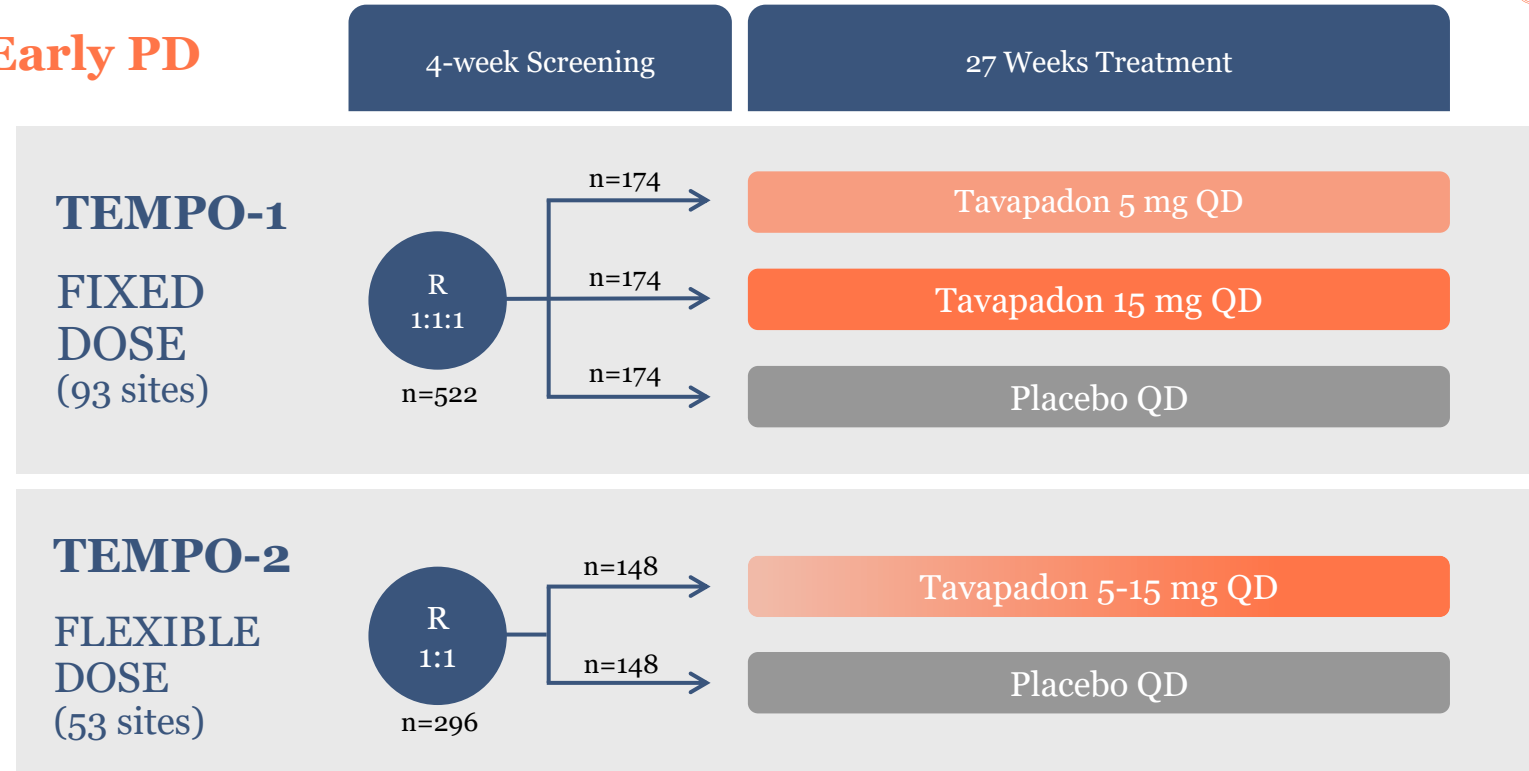
TEMPO-1 & TEMPO-2: Phase 3 in Early PD

Key inclusion criteria

- Adults 40-80 years old
- Baseline MDS-UPDRS⁽¹⁾ Part III Score ≥ 10 and Part II Score ≥ 2
- Modified Hoehn & Yahr⁽²⁾ stage 1 to 2
- No concomitant meds except MAO-B inhibitors

Primary endpoint

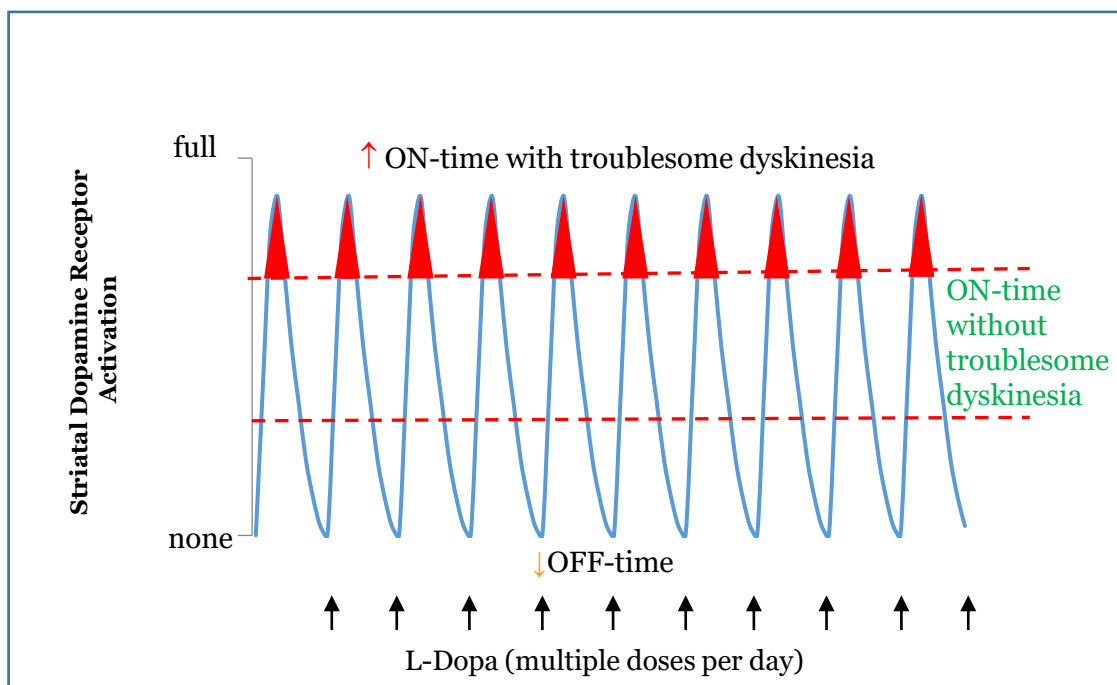
- Change in MDS-UPDRS Parts II+III



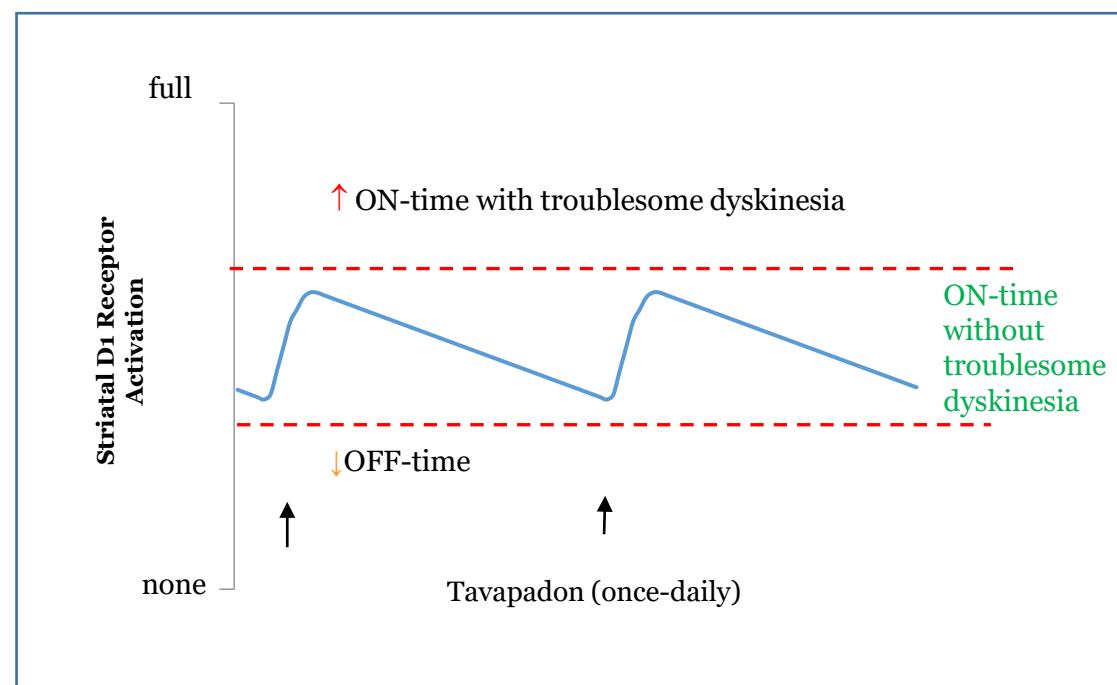
In Late PD, Goal is to Control Motor Fluctuations and Dyskinesias Driven by L-Dopa

L-Dopa vs. Tavapadon in Late-Stage PD¹

L-Dopa is a **FULL** agonist with **SHORT** half-life



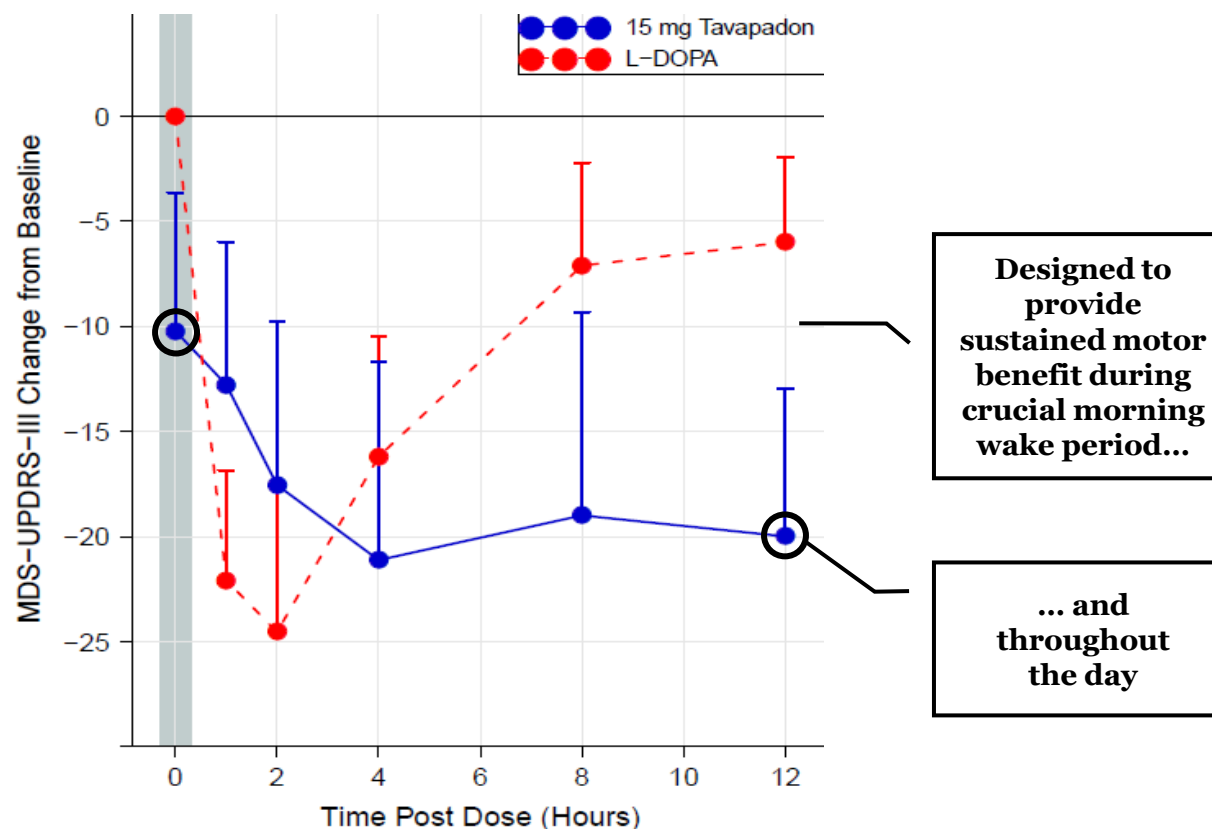
Tavapadon is a **PARTIAL** agonist with **LONG** half-life



30-40% of patients experience dyskinesias within 5 years on L-Dopa²
40% experience OFF-time within 3-5 years on L-Dopa²

Predictable 24-hour Activity → Sustained Effect: Once Daily, Oral Dosing

Study 1005: Tavapadon in Late-Stage PD¹



In an open-label Phase 1b trial, Tavapadon demonstrated motor improvement on par with L-Dopa, sustained due to its 24-hour half-life

Tavapadon TEMPO-3 in Late PD: Data Expected 1H 2023



Registration-directed program includes three global Phase 3 trials optimally designed to maximize treatment effect

TEMPO-3: Phase 3 in Late PD

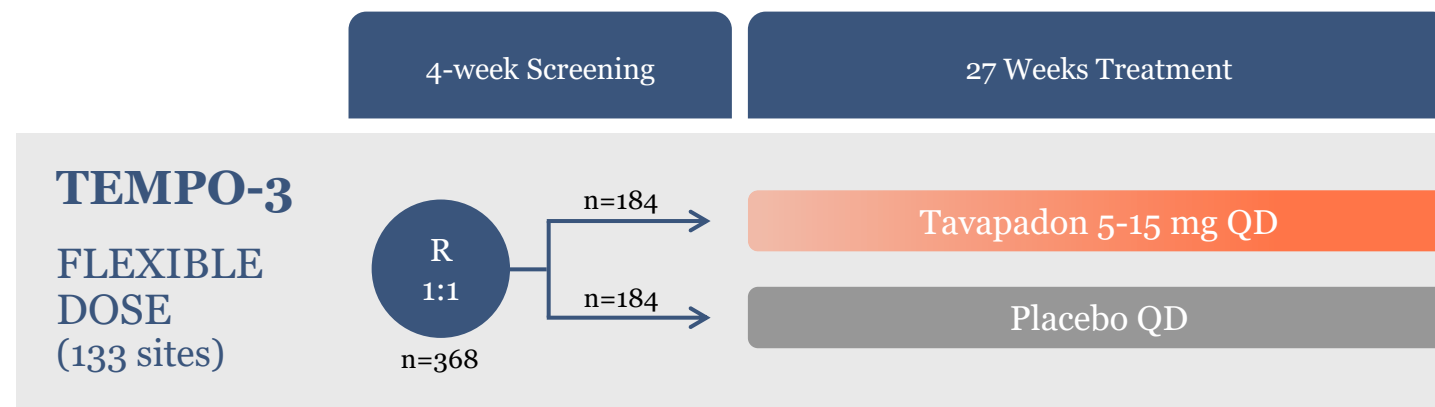
Adjunct to levodopa

Key inclusion criteria

- Adults 40-80 years old
- At least 2.5 hours OFF-time on 2 consecutive days at baseline
- Modified Hoehn & Yahr⁽¹⁾ stage 2 to 3, with response to L-Dopa

Primary endpoint

- Change in ON-time without troublesome dyskinesia



Tavapadon Commercial Potential in Parkinson's

Tavapadon Target Profile



Novel D1/D5 mechanism



Potential similar or better motor control⁽¹⁾



Potential favorable side effect profile⁽²⁾



Once-daily dosing

Pricing & Launch

Branded US price analogs \$8-10K+/year

Payor research supports broad Medicare and Commercial coverage at price of \$8K+ /year

Strong side effect profile and motor control differentiation would reduce reimbursement restrictions

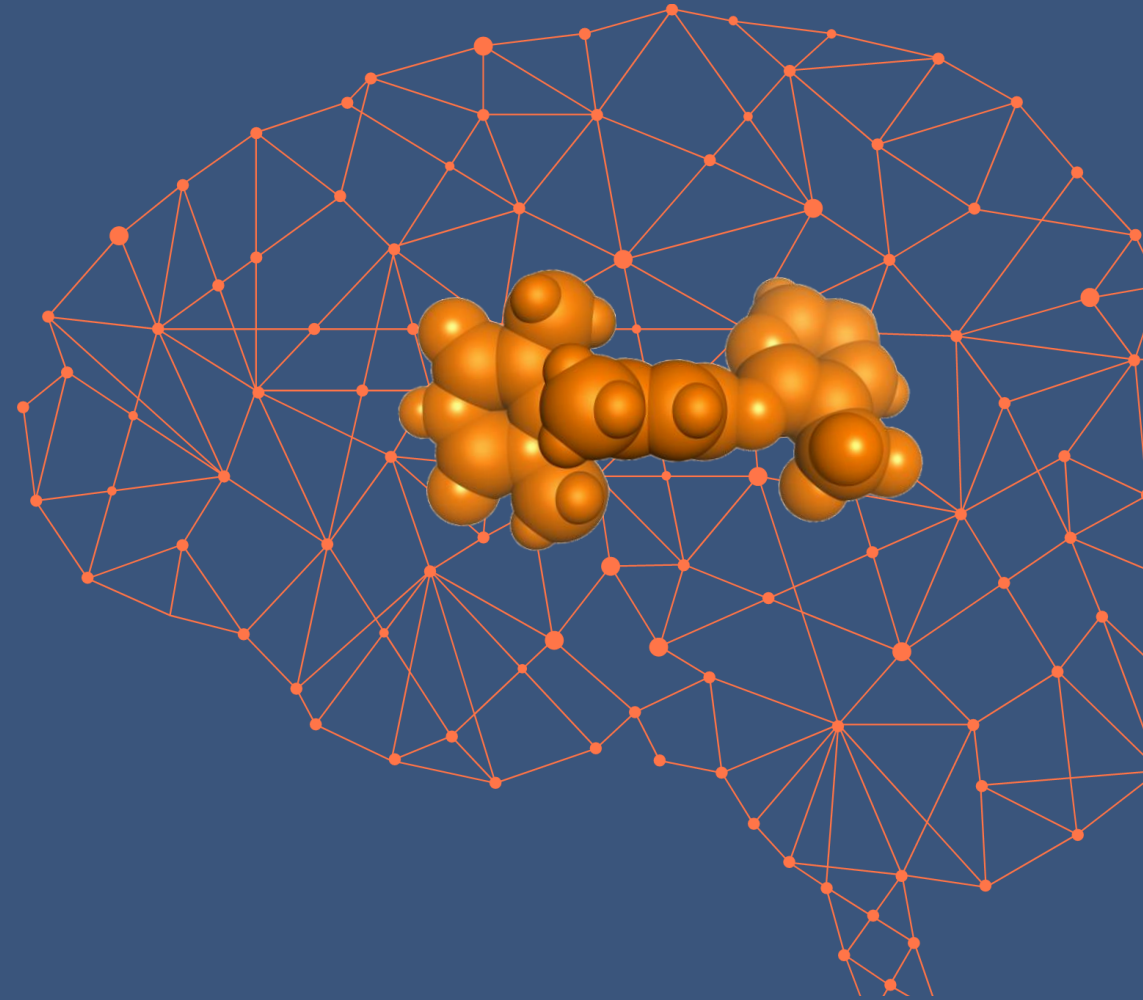
Patients and physician research supports acceptability of branded co-pays for a tavapadon-like differentiated profile



Differentiated profile supports pricing comparable to branded market leaders which have broad reimbursement

CVL-871 in Dementia-Related Apathy

Partial agonist selectively targeting the dopamine D1 receptor with the goal of modulating motivation and reward pathways to address apathy in patients with mild-to-moderate dementia



High Unmet Need in Apathy, which Affects ~50% of Patients with Dementia¹

What is Apathy?

Leading neuropsychiatric symptom in dementia

Social disengagement and loss of emotion leads to:

- Impaired decision-making
- Lack of empathy, affection, or concern
- Loss of interest in personal well-being and relationships
- Inability to initiate and maintain normal daily activities
- Interference with basic function*

CVL-871: Potential to be the First Treatment for Dementia-Related Apathy

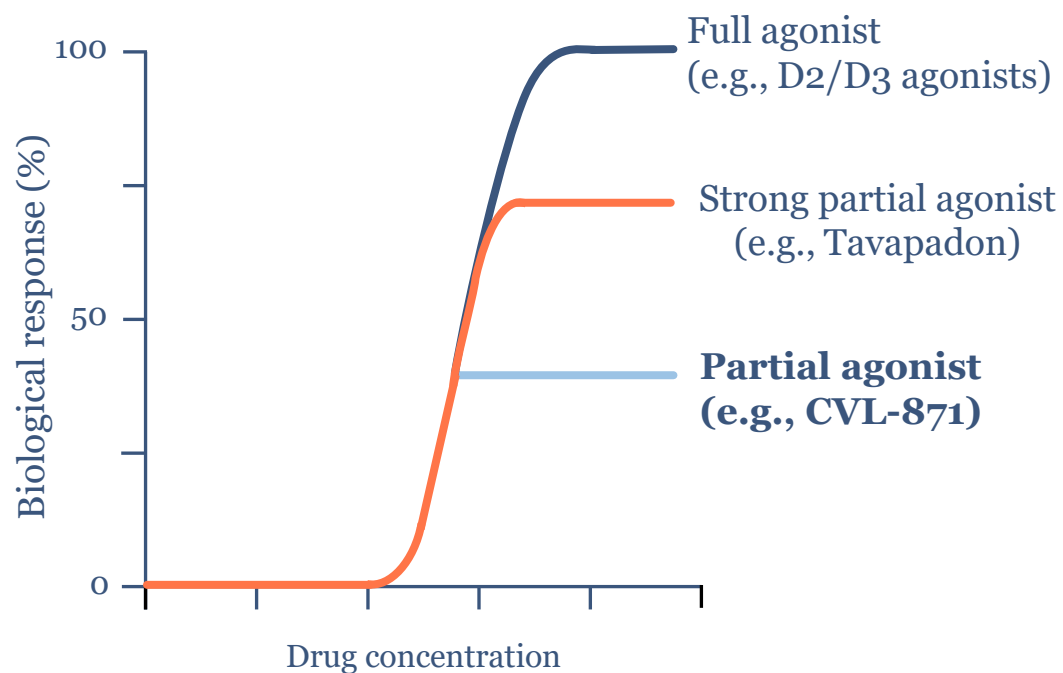
High Unmet Need	>50M Dementia Patients Worldwide ²	>3X Mortality Risk ³	Among strongest predictors of disease progression ⁴	Early institutionalization & ↑ caregiver burden ⁴
No Approved Treatment	Acetylcholinesterase Inhibitors	▷	No proven effect	
Off-label use of...	SSRIs/SNRIs	▷	No established benefit May worsen apathy symptoms	
	Methylphenidate	▷	Schedule II stimulant with CV risk in elderly patients	

CVL-871: D1/D5 Partial Agonism for Dementia-Related Apathy

CVL-871 Summary

- Like tavapadon, CVL-871 is a selective D1/D5 partial agonist
- While tavapadon drives up to ~70% biological response at the D1/D5 receptors, CVL-871 has ~40% partial agonism
- Potentially optimal level of agonism for modulating neuronal pathways related to motivation and reward
- Dopaminergic enhancement may improve apathy based on historical studies of methylphenidate
- Potential non-stimulant option for treatment of dementia-related apathy

Degrees of Agonism (Illustrative)



CVL-871 Phase 2a Exploratory Trial: Data Expected 2H 2022

Phase 2a Trial in Dementia-Related Apathy

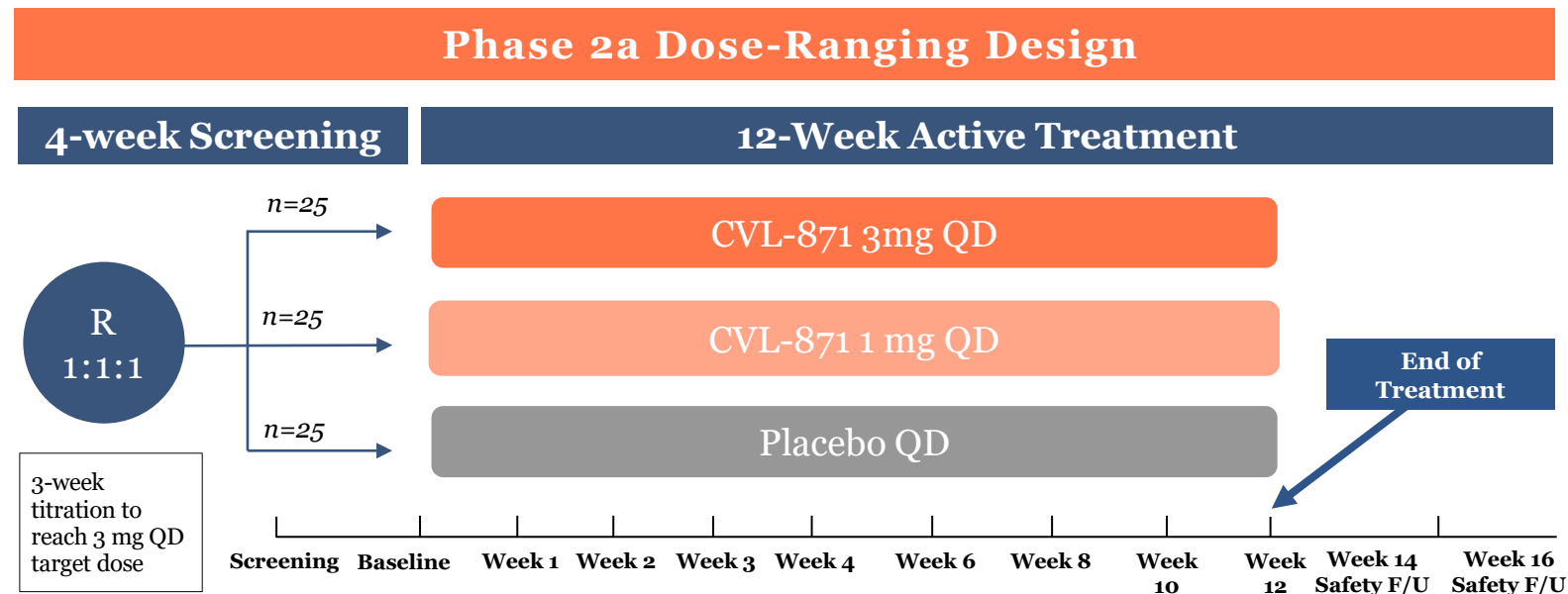
Key inclusion criteria

- Adults 50-85 years old
- NPI-Apathy domain frequency and severity scores each ≥ 2
- Mild-to-moderate dementia
- MMSE 15-26; CDR 0.5-2.0

No primary endpoint

Exploratory efficacy measures

- **Apathy/Global:** NPI/NPI-C, DAIR, AES-C, mADCS-CGIC/CGIS, Caregiver CGIC/CGIS
- **Function:** DAD, Zarit Caregiver Burden
- **Cognition:** ADAS-Cog13, Trail Making A, Digit Span, COWAT



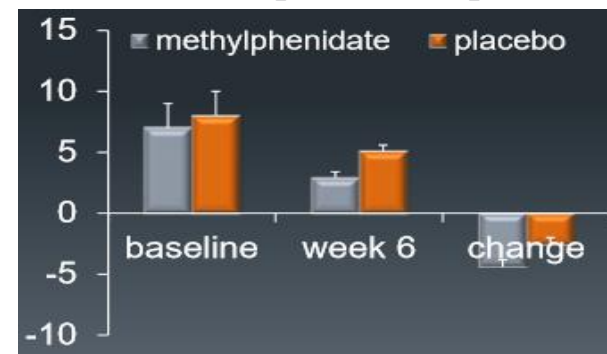
Increased D1 Receptor Activation May Improve Apathy

D1 Activation Potential in Apathy

- Dopamine acting via D1 in the striatum directly promotes motivation and goal-directed behavior
- D1 density reduces with age and reduction in dopamine signaling is associated with behavioral / psychological symptoms of dementia (BPSD)
- Methylphenidate (MPH), an NDRI*, significantly improved apathy in AD patients in 2 independent Phase 2 trials

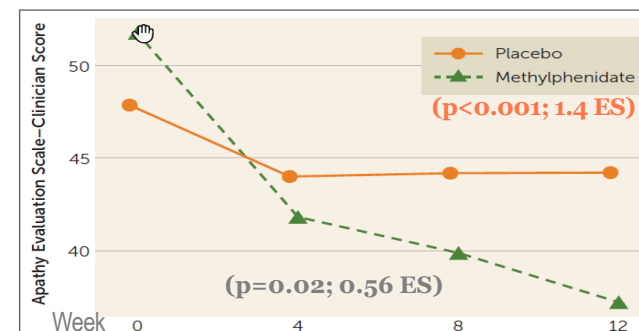
Methylphenidate Phase 2 Trials

ADMET1 Trial: showed NPI Apathy global score improvement of 1.8 points over placebo



ADMET1 Trial-Rosenberg, et al J Clin Psychiatry 2013

Veterans AD Apathy Trial: showed AES-C score improvement of 9.9 points over placebo at week 12

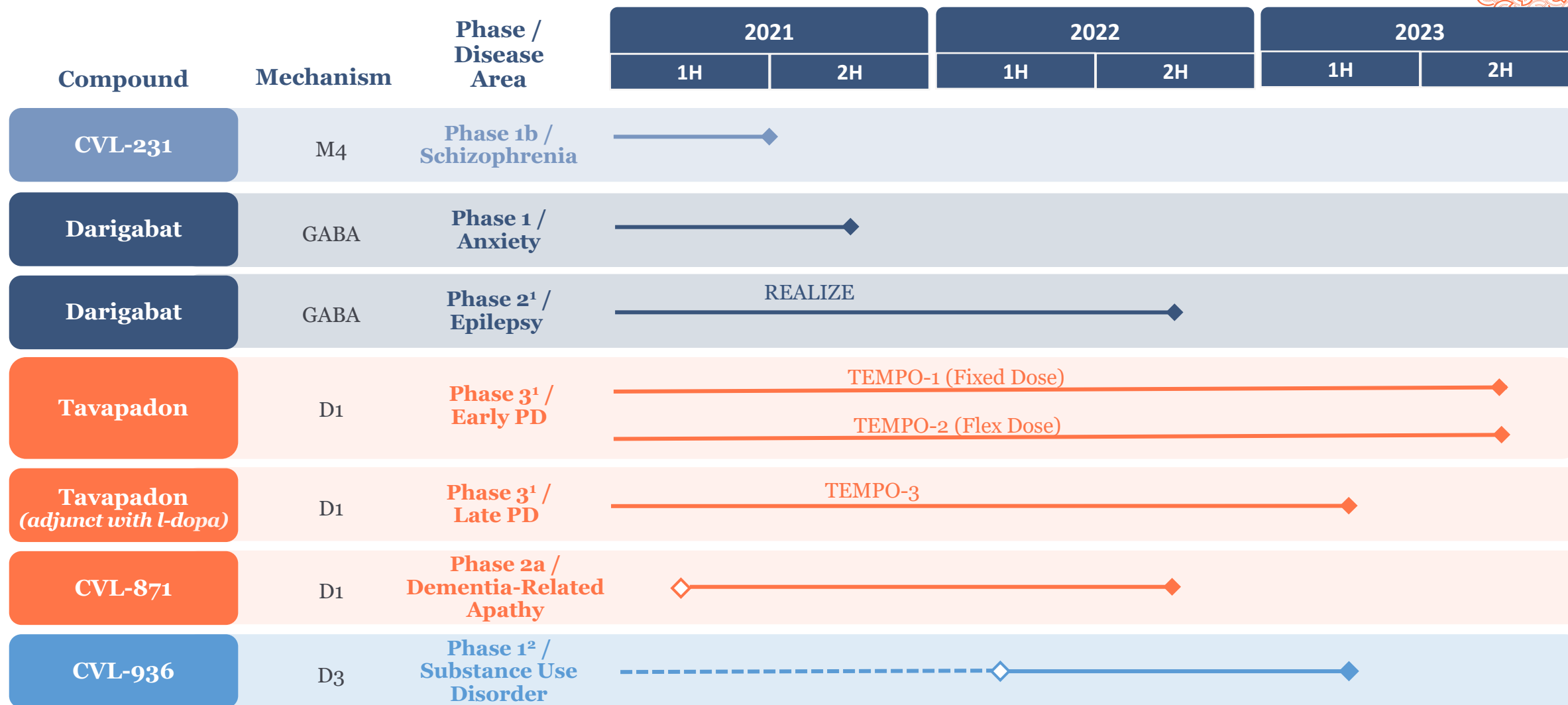


Veterans AD Apathy Trial, Padala et.al, Am J Psychiatry 2018

Transforming the Possible in Neuroscience



Multiple Milestones Expected Over Next Three Years



◇ Estimated Trial Initiation

◆ Estimated Topline Data

Cerevel is Transforming Possibilities for Tomorrow

Multiple Programs Aimed at Providing New Options for Millions of Patients

Tangible near-term value creation

- Schizophrenia
- Epilepsy
- Parkinson's

Expansion to other diseases

- Alzheimer's Psychosis
- Anxiety
- Apathy
- Substance Abuse Disorder

Long-term discovery efforts

Disease-modifying therapies based on human genetics and novel targets addressing:

- Neuronal loss
- Synaptic health

Compound	Mechanism	Disease Area	Discovery	Pre-clinical	Phase 1	Phase 2	Phase 3
CVL-231	M4 PAM	Schizophrenia					
Darigabat	GABA _A α2/3/5 PAM	Epilepsy					
Tavapadon	D1/D5 Partial Agonist	Early Parkinson's					
Tavapadon (adjunct with L-Dopa)	D1/D5 Partial Agonist	Late Parkinson's					

Compound	Mechanism	Disease Area	Discovery	Pre-clinical	Phase 1	Phase 2	Phase 3
CVL-231	M4 PAM	Schizophrenia					
Darigabat	GABA _A α2/3/5 PAM	Epilepsy					
Darigabat	GABA _A α2/3/5 PAM	Anxiety					
Tavapadon	D1/D5 Partial Agonist	Early Parkinson's					
Tavapadon (adjunct with L-Dopa)	D1/D5 Partial Agonist	Late Parkinson's					
CVL-871	D1/D5 Partial Agonist	Dementia-Related Apathy					
CVL-936	D3- Preferring Antagonist	Substance Use Disorder					

Compound	Mechanism	Disease Area	Discovery	Pre-clinical	Phase 1	Phase 2	Phase 3
CVL-231	M4 PAM	Schizophrenia					
CVL-865	GABA _A α2/3/5 PAM	Epilepsy					
CVL-865	GABA _A α2/3/5 PAM	Anxiety					
Tavapadon	D1/D5 Partial Agonist	Early Parkinson's					
Tavapadon (adjunct with L-Dopa)	D1/D5 Partial Agonist	Late Parkinson's					
CVL-871	D1/D5 Partial Agonist	Dementia-Related Apathy					
CVL-936	D3- Preferring Antagonist	Substance Use Disorder					
CVL-354	KORA	MDD / SUD					
Lead Optimization	PDE4B	MDD / Schizophrenia					
Lead Optimization	M4 Agonist	Schizophrenia					
Lead Optimization	LRRK2	Parkinson's					

50M+ Patients WW



100M+ Patients WW



Premier Neuroscience Company

Appendix



Who we are is in our name

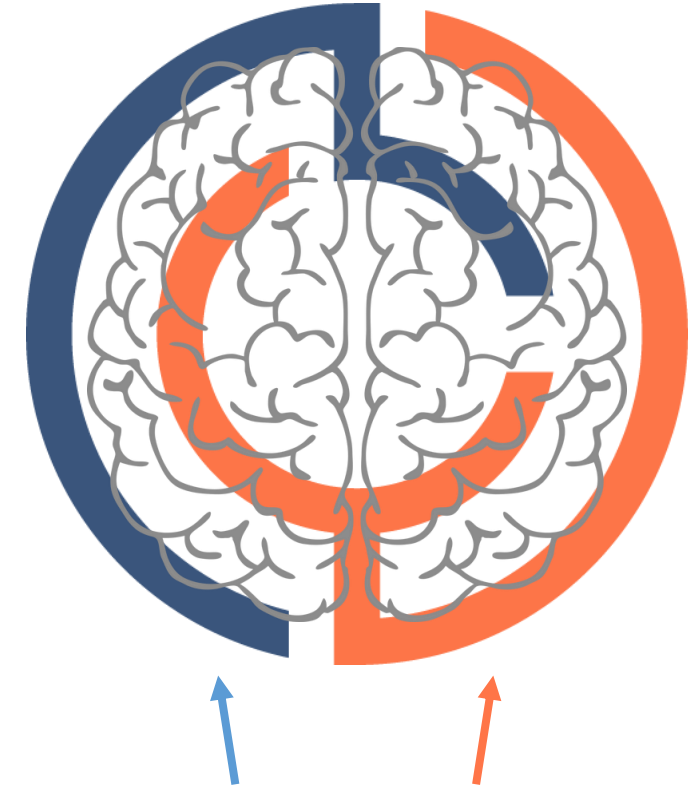


cere = cerebrum

revel = revelation/reveal

We are bold thinkers, deep experts, resilient pathfinders, and transparent partners who push the boundaries of scientific understanding to unlock breakthrough CNS therapies that could have real impact on people's lives.

Brain hemispheres



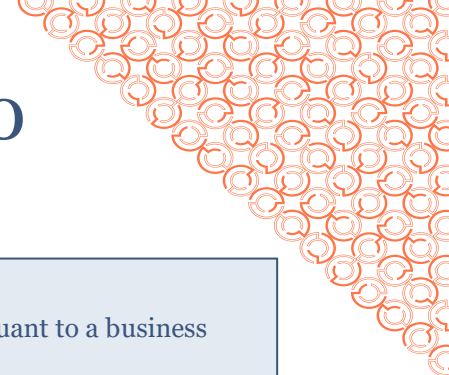
Lock in key

Selective / Targeted Mechanisms

Cerevel Clinical Pipeline: Broad, Deep and Diverse

Compound	Disease Area	Discovery	Pre-clinical	Phase 1	Phase 2	Phase 3	Upcoming Milestone	Mechanism
Tavapadon	Early Parkinson's						Ph. 3 Data 2H 2023	D1/D5 Strong Partial Agonist ★
Tavapadon (adjunct with L-Dopa)	Late Parkinson's						Ph. 3 Data 1H 2023	
Darigabat	Epilepsy						Ph. 2 Data 2H 2022	GABA _A α2/3/5 PAM ★
Darigabat	Anxiety						Ph. 1 Data 2H 2021	
CVL-231	Schizophrenia						Ph. 1b Data Mid-year 2021	M4 PAM ★
CVL-871	Dementia-Related Apathy						Ph. 2a Data 2H 2022	D1/D5 Partial Agonist ★
CVL-936	Substance Use Disorder						Under evaluation	D3 Preferring Antagonist

Combination with ARYA II Completed on October 27, 2020



Transaction Summary

- Cerevel Therapeutics, Inc. (“Cerevel”) and ARYA Sciences Acquisition Corp II (“ARYA II”, Nasdaq: ARYB) merged pursuant to a business combination agreement between ARYA II and Cerevel
 - Cerevel is a clinical-stage biopharmaceutical company that combines a deep understanding of the biology and neurocircuitry of the brain with advanced chemistry and central nervous system (CNS) receptor pharmacology to discover and develop new therapies
 - ARYA II was a special purpose acquisition company sponsored by Perceptive Advisors
- Transaction closed on 10/27/20, Cerevel Therapeutics Holdings, Inc. now trading on Nasdaq as ticker **CERE**

Premier Specialist Investor Base

- Provided Cerevel with premier investor base and resources to continue executing on its development plan.
- Shareholders of the combined company include current Cerevel and ARYA II shareholders as well as top-tier biotech / life sciences investors, including Perceptive

Use of Proceeds

- Net proceeds of \$440M
 - Proceeds expected to fund Cerevel's R&D programs, including M4 PAM (CVL-231) in schizophrenia, darigabat in anxiety and epilepsy, D1 partial agonist (tavapadon) in Parkinson's and earlier-stage clinical programs
 - Expected to provide runway into 2023

Key Management and Board

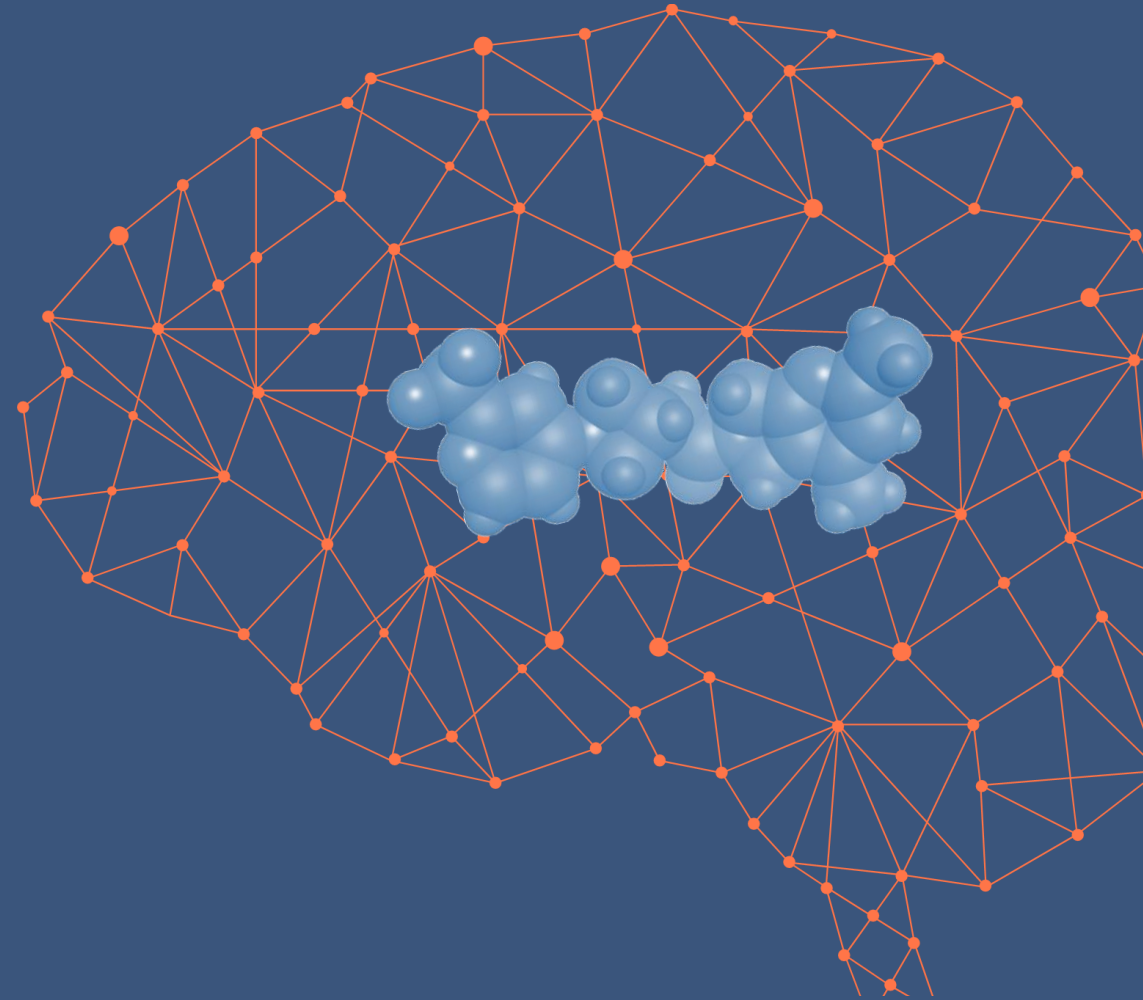
- Combined company led by Cerevel Chief Executive Officer & Chairperson, Tony Coles, M.D.

Combination with ARYA II – Key Highlights

- Provides a faster path to becoming a public company
 - Addresses one of the leading questions from potential crossover investors by enabling Cerevel to go public in one step vs. a typical two step process including crossover and IPO
 - Provides an investment structure for public investors to enable a potential business combination that appropriately capitalizes Cerevel while meaningfully reducing market risk
- Capitalizes Cerevel with \$440 million⁽¹⁾ raise through the reverse merger and PIPE to fund broad portfolio of neuroscience assets
 - Expected to provide cash runway for key catalysts into 2023, including:
 - Up to six data readouts across diversified pipeline of early and late stage programs
 - Additional IND filings for novel MOAs in new indications
- Price discovery streamlined and reduced execution risk in volatile markets
 - Satisfies investors' desire for larger capital raise to meet increased market demand
- Ability to establish premier shareholder base capable of supporting the company into the future
- Establish a broad syndicate of banks and research analysts that follow the stock post closing

M4 PAM (CVL-231) in Schizophrenia

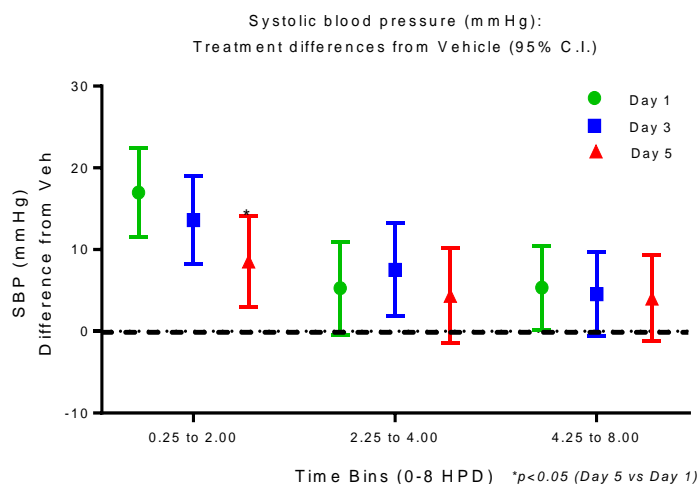
Additional Slides



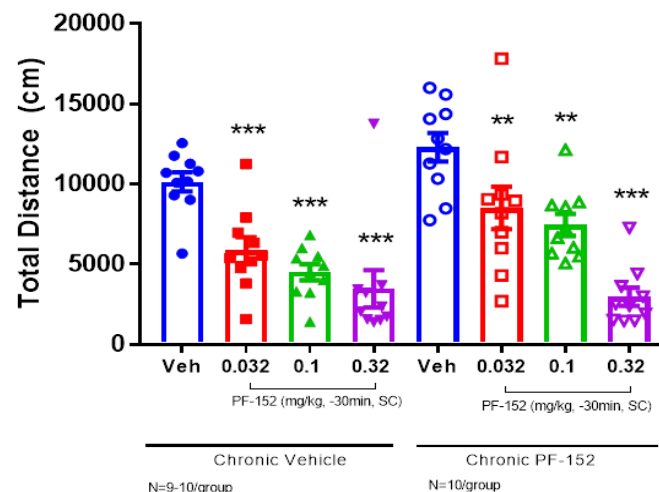
Cardiovascular Effects may be Attenuated with Titration and Repeat Dosing

Repeated dosing of M4 Agonist Tool in rodents showed attenuation of cardiovascular effects without impact on antipsychotic activity; in addition, CVL-231 showed attenuation of heart effects in a 3-month canine toxicology study

5-Day repeat dosing of M4 Agonist Tool: Attenuation of Blood Pressure Effects in Mice



14-Day repeat dosing of M4 Agonist Tool: No Attenuation of Antipsychosis Effects in Mice



3-Month Study of CVL-231: Attenuation of Heart Rate Effects in Canines

- On Day 1, observed heart rate increases were statistically significant and outside normal range
- On Days 43 and 90, heart rate increases were small and not statistically significant; all mean heart rate values were within normal range and not considered adverse



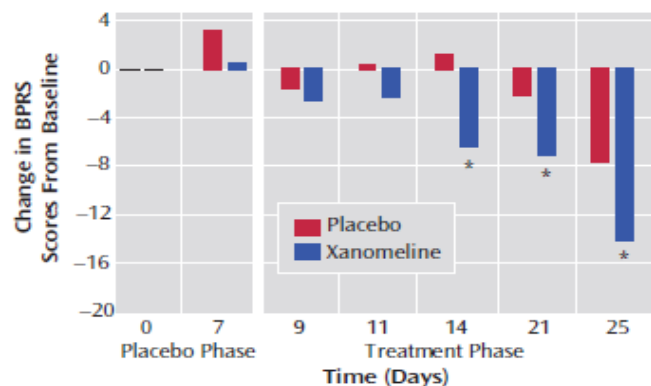
Clinical translation: KarXT showed an average increase in resting heart rate of 5.5 beats per minute with a downward trend after the second week

Xanomeline Clinical Data: Compelling Activity, Limited by Side Effect Profile

Xanomeline (Non-selective Agonist) Impacted Symptoms...

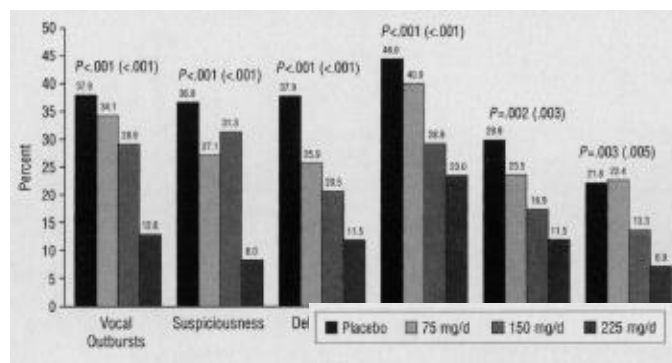
2008 Phase 2 in Schizophrenia

Statistically significant impact on **total BPRS** and **PANSS** scores in schizophrenia patients¹



1997 Phase 2 in Alzheimer's Disease

Statistically significant impact on **agitation** and **other psychosis-related endpoints** in Alzheimer's patients²



...But Development Was Limited by GI Side Effects

Table 3. Adverse Events*

Event	Placebo (n=87)	Dose†			Total (N=342)	P‡
		Low (n=85)	Medium (n=83)	High (n=87)		
Sweating	4 (4.6)	12 (14.1)	38 (45.8)	66 (75.9)	120 (35.1)	<.001
Nausea	17 (19.5)	24 (28.2)	29 (34.9)	45 (51.7)	115 (33.6)	<.001
Vomiting	8 (9.2)	11 (12.9)	33 (39.8)	37 (42.5)	89 (26.0)	<.001
Dyspepsia	7 (8.0)	20 (23.5)	23 (27.7)	21 (24.1)	71 (20.8)	.007
Chills	1 (1.1)	8 (9.4)	22 (26.5)	32 (36.8)	63 (18.4)	<.001
Chest pain	1 (1.1)	5 (5.9)	13 (15.7)	10 (11.5)	29 (8.5)	.004
Increased salivation	0 (0)	2 (2.4)	6 (7.2)	21 (24.1)	29 (8.5)	<.001
Syncope	4 (4.6)	3 (3.5)	11 (13.3)	11 (12.6)	29 (8.5)	.03
Fecal incontinence	0 (0)	4 (4.7)	1 (1.2)	5 (5.8)	11 (3.2)	.04
Nausea and vomiting	2 (2.3)	0 (0)	1 (1.2)	7 (8.0)	10 (2.9)	.009
Dysphagia	1 (1.1)	0 (0)	2 (2.4)	6 (6.9)	9 (2.6)	.03

*Only events statistically significant at $P < .05$ are given. Values are number (percentage) of patients unless otherwise indicated.
 †Low-dose group received 25 mg of xanomeline tartrate 3 times a day; medium, 50 mg 3 times a day; high, 75 mg 3 times a day.
 ‡Pearson χ^2 test.

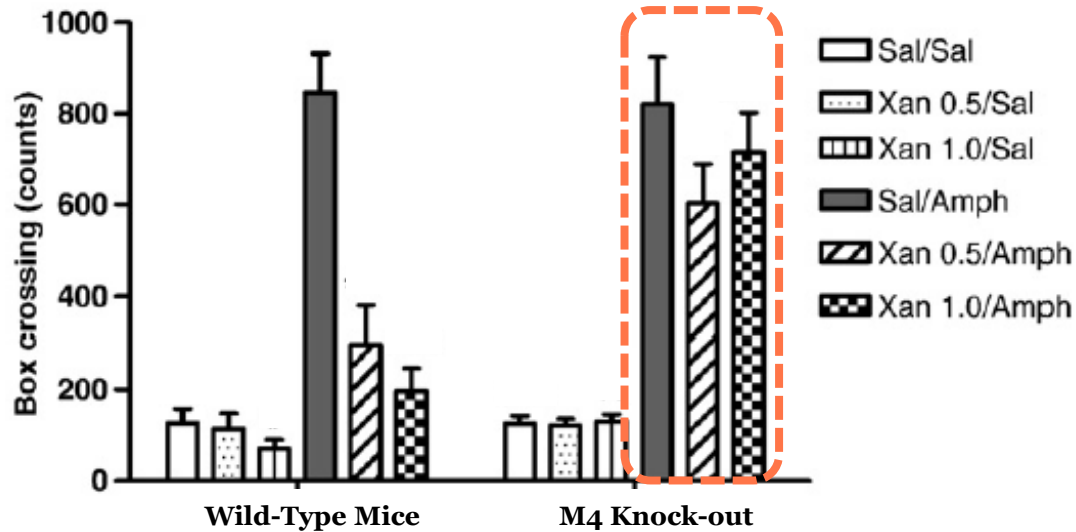
Xanomeline's high rate of nausea, vomiting and dyspepsia is likely driven by non-selective muscarinic agonism

Mechanism Supported by Phase 2 Data for KarXT

- Karuna is developing a BID fixed-dose combination of xanomeline with trospium to offset side effects of GI, dry-mouth and constipation
- In a 5-week Phase 2 study, KarXT demonstrated an 11.6 point reduction in PANSS total score from baseline vs. placebo ($p < 0.0001$)
- ~70 completers in each arm, with discontinuation rates similar between placebo and treatment group
- Represents a robust reproduction of 2008 xanomeline Phase 2 data in schizophrenia

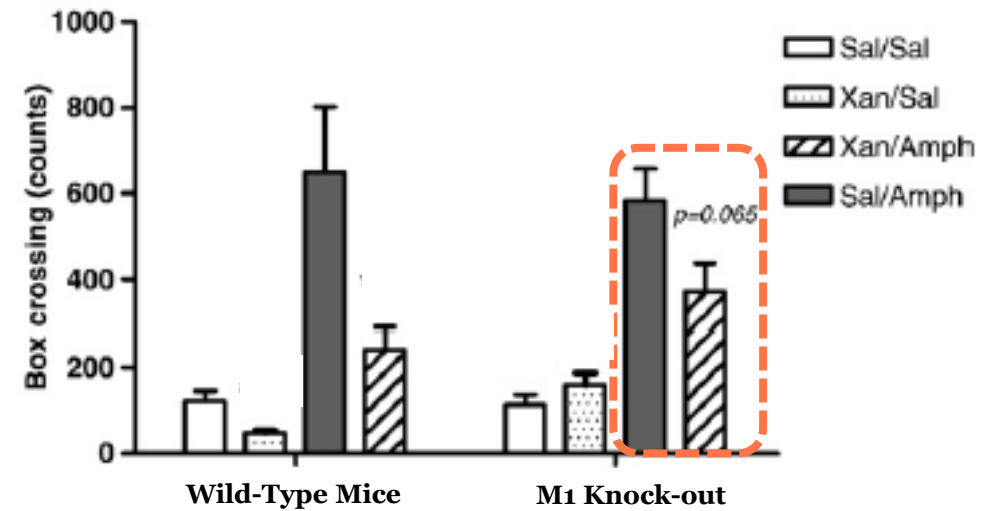
Preclinical Evidence: M4 Modulation Drives Antipsychosis

No Effect of Xanomeline in M4 Knock-out Mouse Model



Xanomeline had no effect on amphetamine-induced hyperactivity in M4 knock-out mice

Reduction in Hyperactivity in M1 Knock-out Mice



Xanomeline reduced hyperactivity in M1 knock-out mice



In mouse studies, M4 receptors drive the antipsychotic activity of xanomeline

Important Insights on Side Effects of M4 PAM

Results of a Phase 1 SAD trial indicated CVL-231 was generally well-tolerated with asymptomatic transient effects on heart rate and blood pressure

Phase 1 SAD Trial (N=17)

Tested doses up to 30 mg

Relatively well tolerated with no SAEs

Most frequently reported treatment-emergent AEs: fatigue, dizziness, headache and dry mouth

Moderate treatment-emergent transient increases in blood pressure and pulse rate observed

Cardiovascular effects were asymptomatic and transient in nature

Insights

Preclinical studies show CV effects attenuated with repeat dosing

KarXT data also suggest that CV effects attenuate over time with repeat dosing

Tolerability may be differentiated in schizophrenia patients and CV effects may be attenuated with repeat dosing and/or titration.

Phase 2 data for MK-7622 (M1 PAM) in Alzheimer’s disease

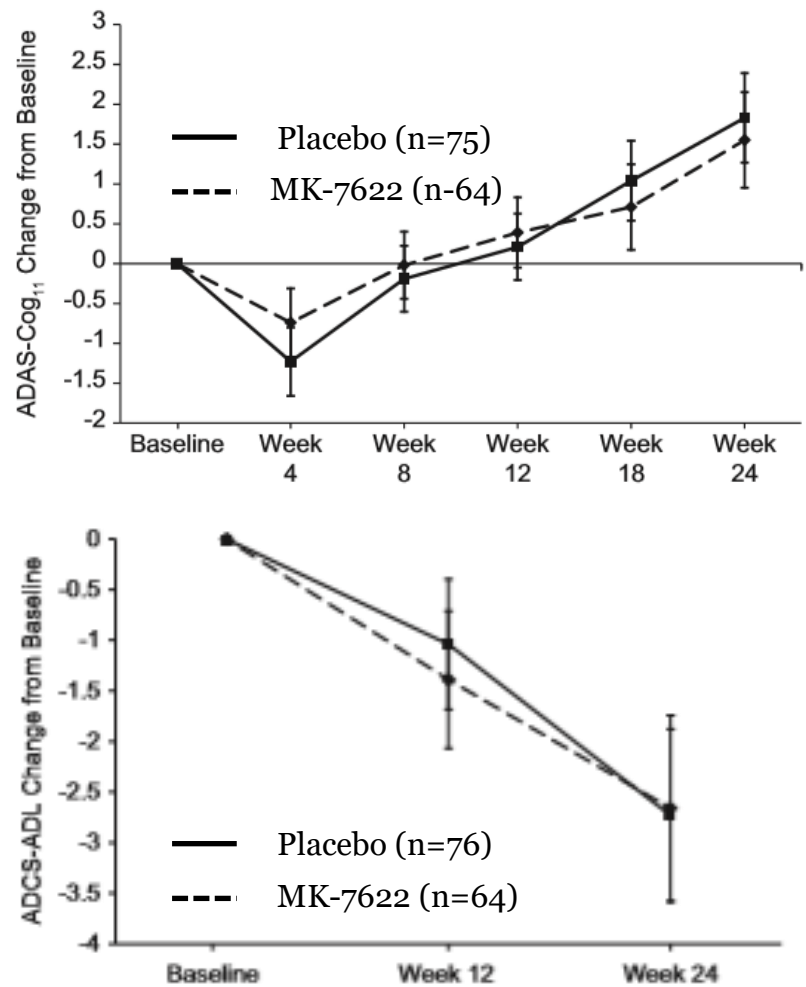
Summary

- Randomized double-blind proof-of-concept trial as adjunctive therapy in mild-to-moderate Alzheimer’s disease
- Conducted by Merck; data published 2018
- Trial stopped early for futility

Results

- **No difference from placebo on either cognition or activities of daily living (ADL) scales**
- Discontinuation rate of 16% on MK-7622 vs 6% on placebo
- Cholinergically-related adverse event rate of 21% on drug vs 8% on placebo

Results in Cognition and ADL

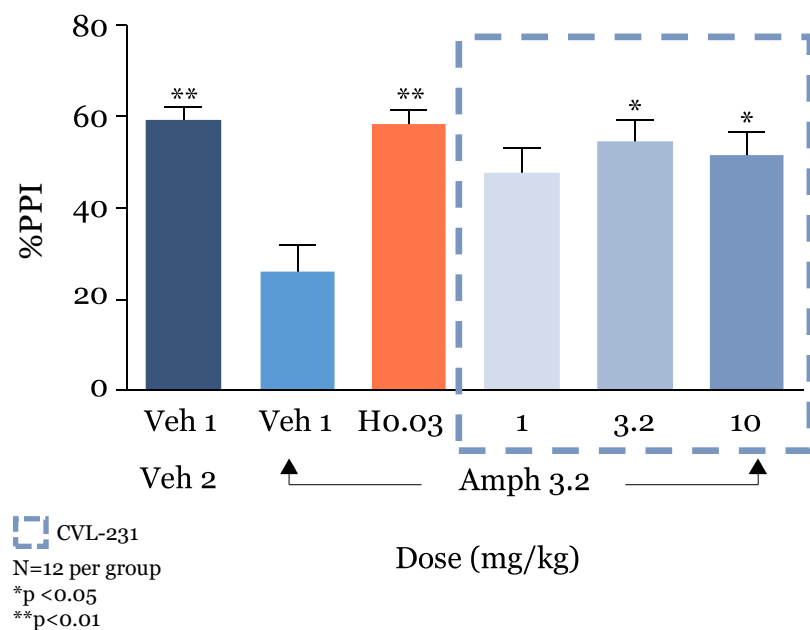


Side Effect Profile

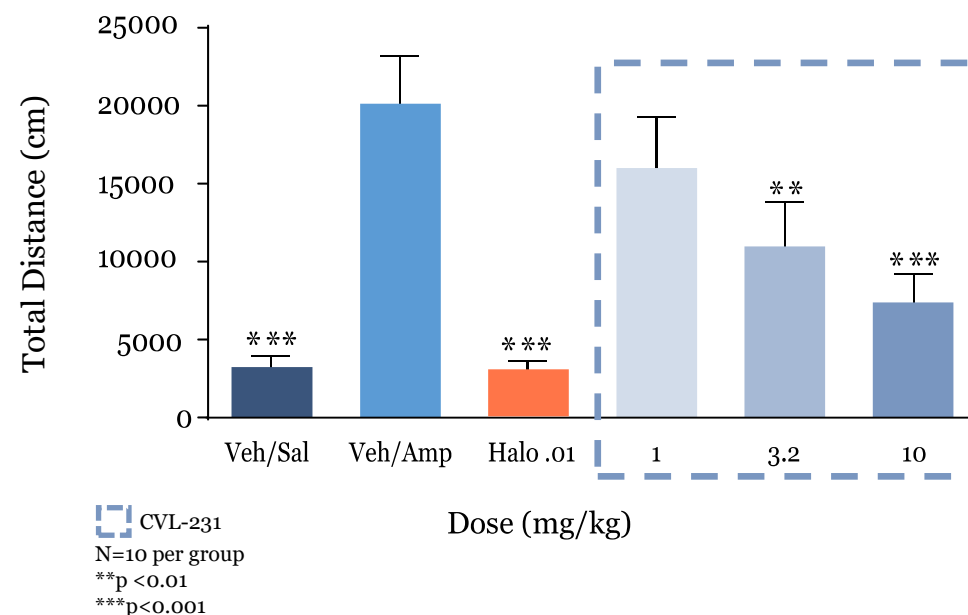
Most Common AEs (>5%)	MK-7662 (n=119)	Placebo (n=120)
Diarrhea	18 (15.1%)	7 (5.8%)
Headache	11 (9.2%)	6 (5.0%)
Rhinorrhea	7 (5.9%)	1 (0.8%)
Urinary Incontinence	6 (5.0%)	0 (0.0%)
Weight Decrease	6 (5.0%)	2 (1.7%)
Urinary Tract Infection	6 (5.0%)	7 (5.8%)
Fall	2 (1.7%)	6 (5.0%)

M4 PAM Preclinical Data in Psychosis

CVL-231 showed similar effect to haloperidol in reversing amphetamine-disrupted Pre-pulse Inhibition (PPI) in rats



CVL-231 showed dose-dependent reductions on amphetamine-induced locomotion in rats



In multiple rodent models of psychosis, CVL-231 demonstrated antipsychotic activity consistent with atypical antipsychotics

Potential Indications for M4 PAM Beyond Schizophrenia

Pipeline in a Pill

**Goal to be a novel MOA
and next generation treatment
in Schizophrenia**

Aiming for a Side Effect and Tolerability
Profile Appropriate for Chronic Use in
Elderly Populations

Potential Large Indications Worldwide

▶	Schizophrenia	~21M Patients
▷	Alzheimer's Psychosis	~20M Patients
▷	Cognition	>50M Patients
▷	PD-L1D	~5M Patients



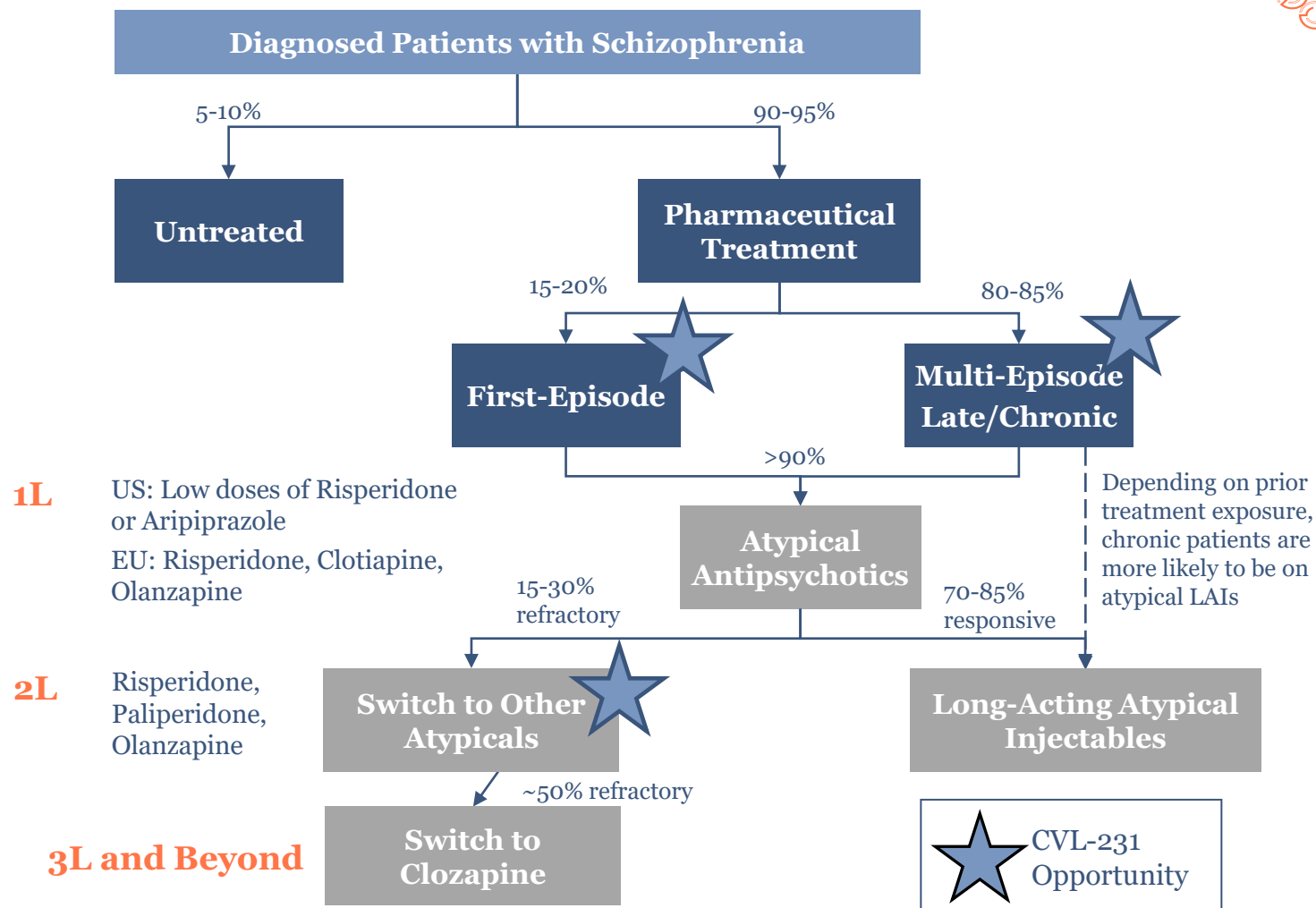
Potential to expand use outside of core schizophrenia population
to behavioral and psychological symptoms of dementia

CVL-231 Commercial Potential in Schizophrenia

Multiple Potential Entry Points for CVL-231 in the Treatment Paradigm

Potential for CVL-231 to be a New Standard of Care

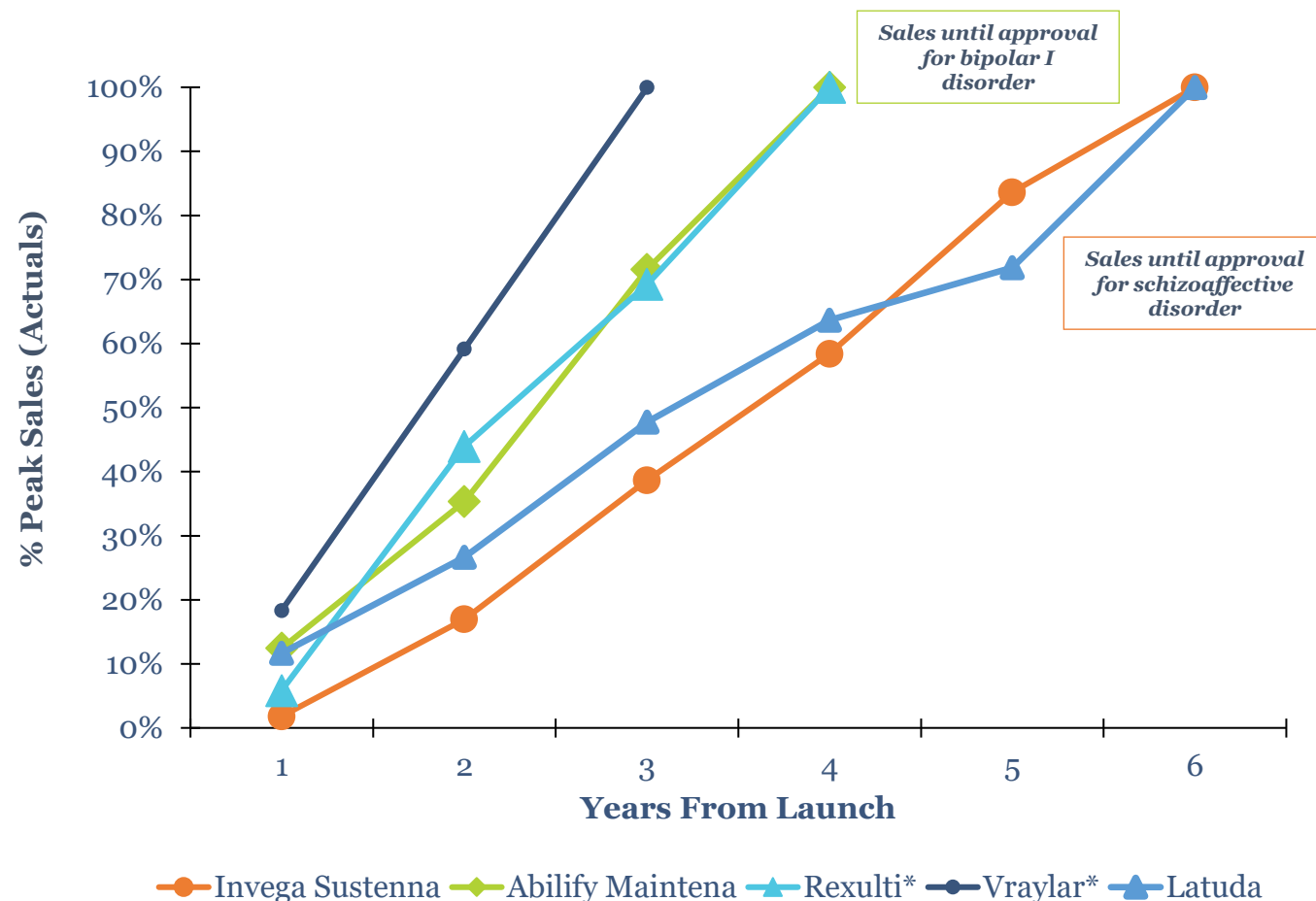
- Potential for improved SE profile could position CVL-231 as first-line treatment of newly diagnosed and ongoing schizophrenia patients, including DRP
- If an improved tolerability and metabolic profile is demonstrated, CVL-231 could displace atypical antipsychotics in patients with treatment-related side effects



Schizophrenia Therapies: Rapid Historic Uptake Despite Limited Differentiation

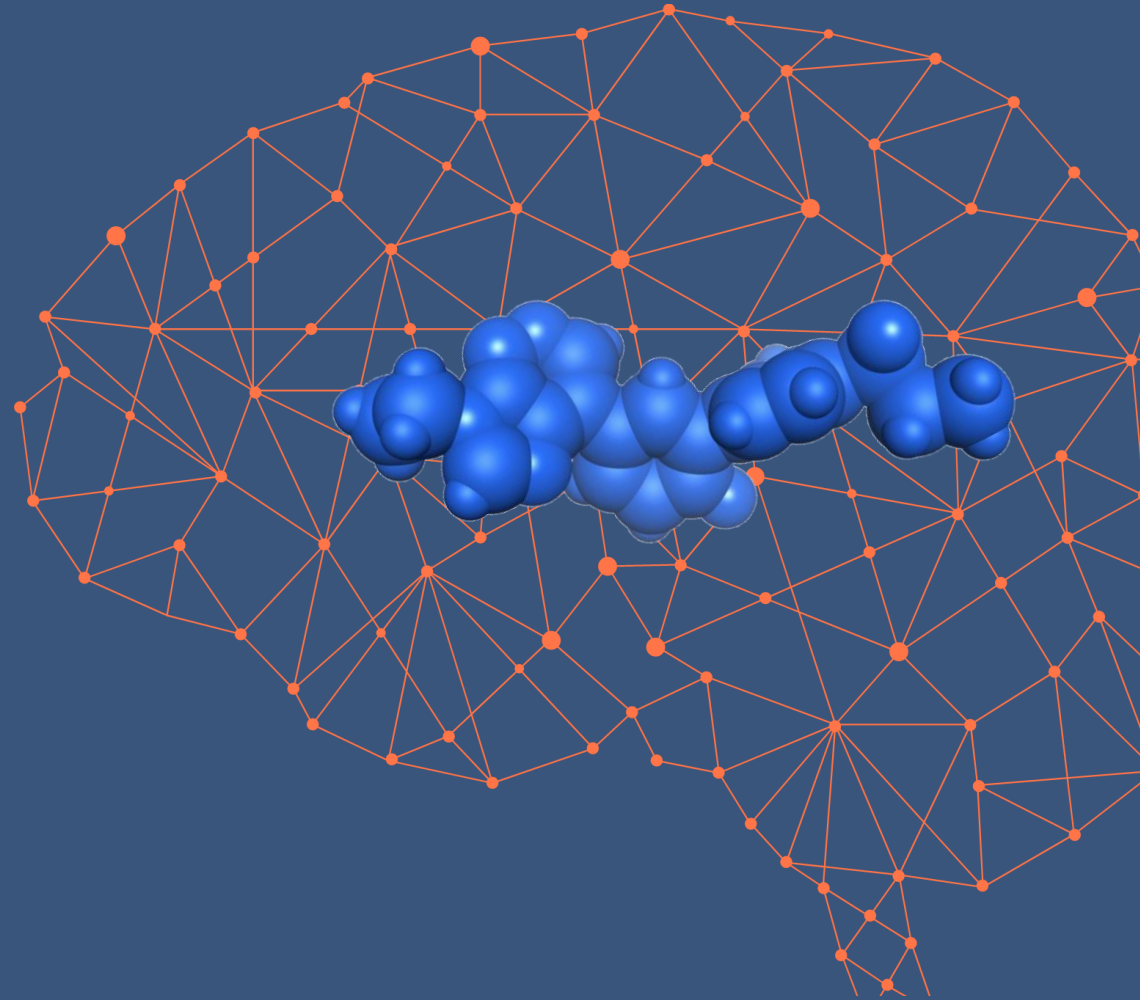
Drug	US 2018 Schizophrenia Sales	2018 US Share
Latuda (lurasidone)	\$973M	13.5%
Invega Sustenna (paliperidone LAI)	\$981M	6.2%
Rexulti (brexpiprazole)	\$449M	8.1%
Abilify Maintena (aripiprazole LAI)	\$331M	2.1%
Vraylar (cariprazine)	\$164M	2.6%

Schizophrenia US Sales Ramp – Actuals
(through 2018 or until first non-schizophrenia indication launch)

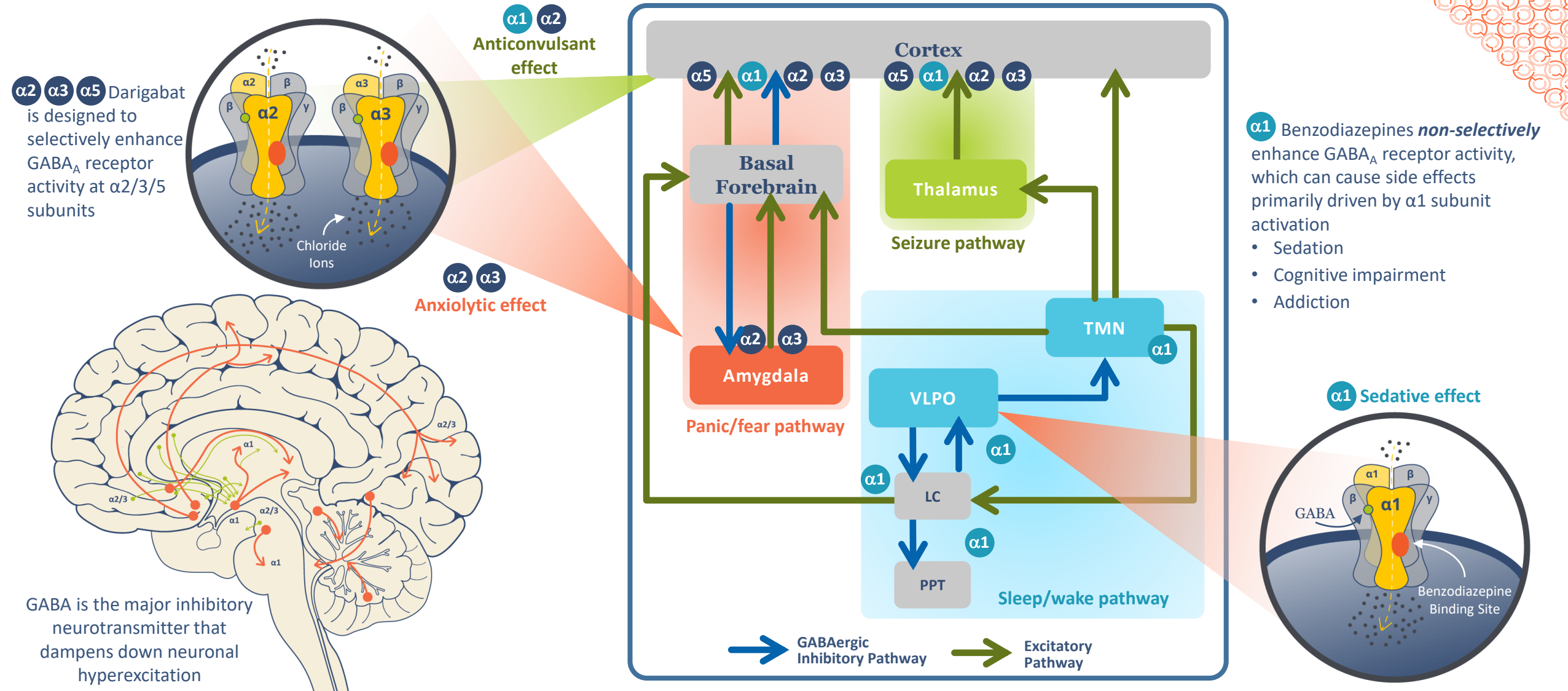


Darigabat in Epilepsy

Additional Slides

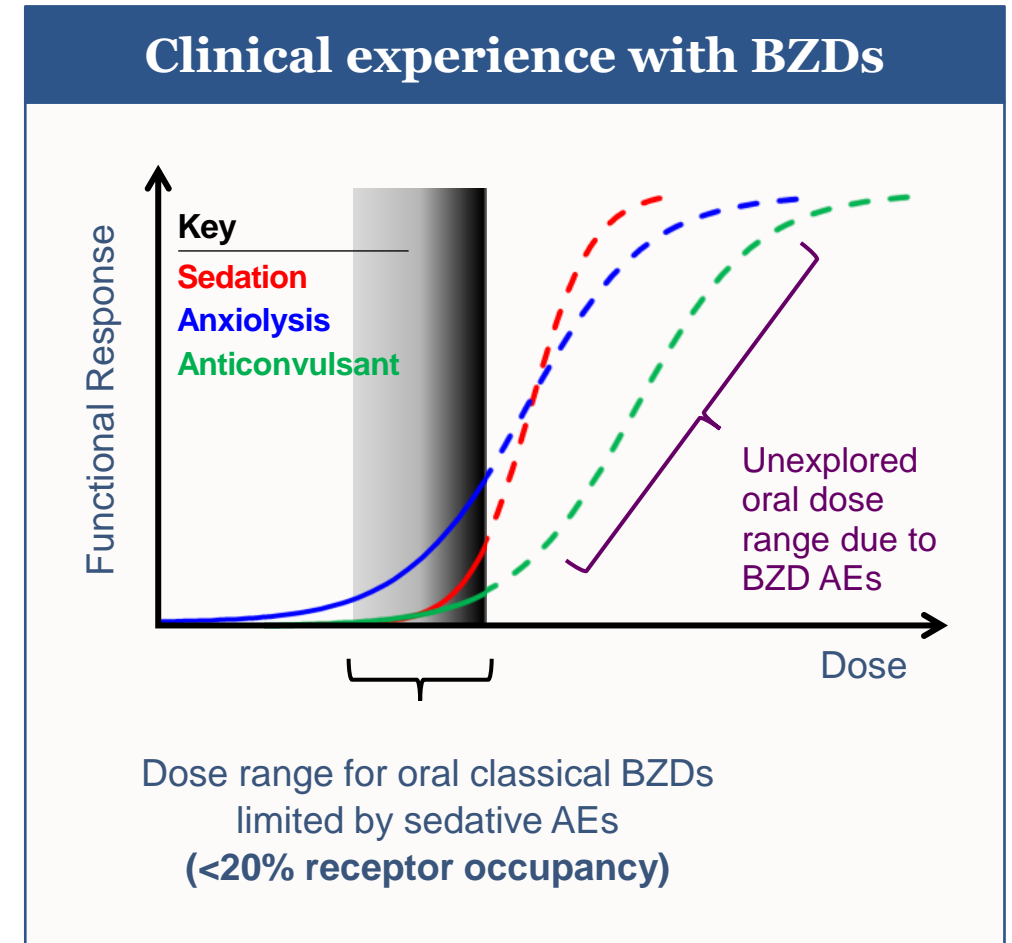


Darigabat Mechanism: Selective $\alpha 2/3/5$ GABA_A Receptor PAM



The Problem With Benzodiazepines (abridged...)

- BZDs are efficacious in a range of indications but use and dose is limited by adverse events, even at low receptor occupancy
 - Sedation, somnolence, cognitive impairment, falls, overuse, misuse and addiction
- In general, BZDs are used acutely in epilepsy but not indicated for chronic use due to tolerance or loss of efficacy
- Darigabat has the potential to be used chronically by minimizing adverse events, risk of tolerance and abuse

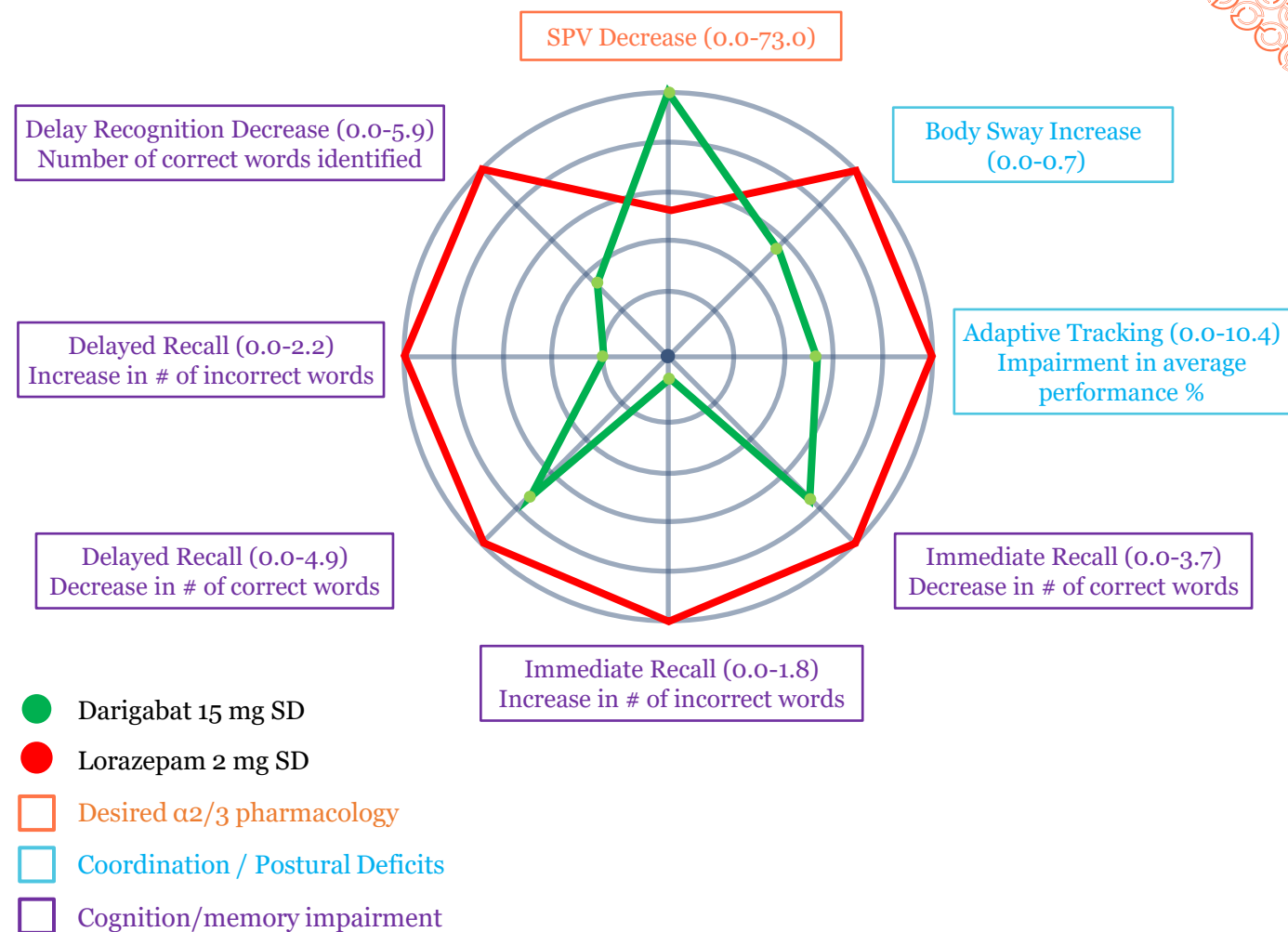
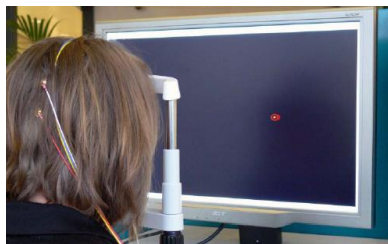


Darigabat: Favorable Pharmacology in NeuroCart, Differentiated From a BZD

NeuroCart is a comprehensive battery of tests to evaluate CNS functional domains

Darigabat first-in-human study tested the following brain functions based on known GABA_A receptor pharmacology:

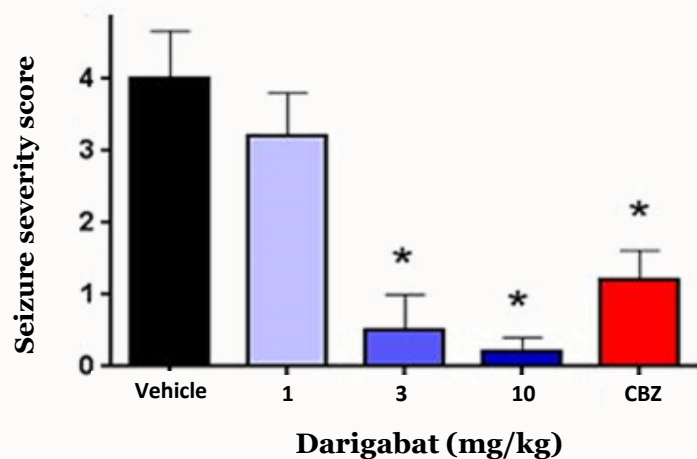
- Saccadic peak velocity (SPV) - desired $\alpha 2/3$ pharmacology
- Body sway - undesired $\alpha 1$ pharmacology
- Adaptive tracking - undesired $\alpha 1$ pharmacology
- Visual-verbal learning test - undesired $\alpha 1/5$ pharmacology
- Relative to 2 mg lorazepam, darigabat demonstrated a larger decrease in SPV and smaller impairment on body sway, adaptive tracking and cognitive tests



Darigabat is Anticonvulsant in a Range of Preclinical Models

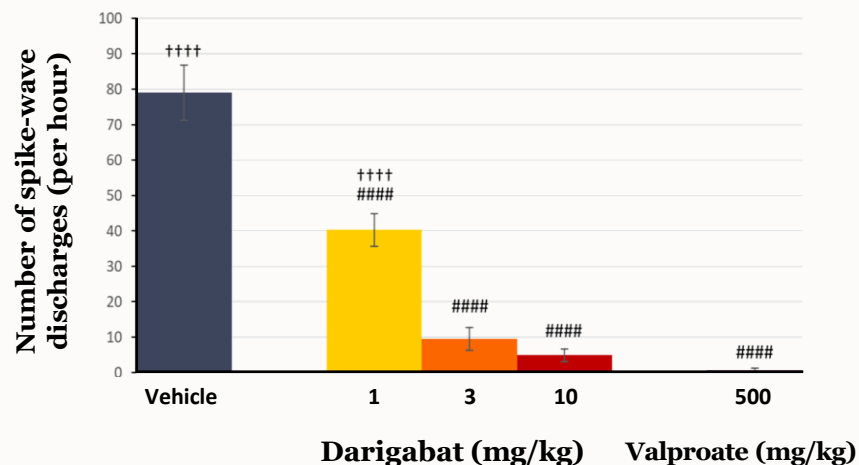
- Strong correlation of animal models of seizures translating to clinical activity across mechanism
- Darigabat demonstrated broad spectrum activity at $\sim >50\%$ receptor occupancy
 - Darigabat is active in pentylenetetrazol-induced seizures
 - Amygdala kindling is a validated model for predicting activity in focal seizures
 - Genetic absence epilepsy rat model predictive of activity in absence (generalized) seizures

Amygdala Kindled Seizures in Rats: Focal Seizures



* p<0.05

Genetic Absence Epilepsy in Rats: Generalized Seizures



†††† p<0.0001 vs. valproate; #### p<0.0001 vs. vehicle

Darigabat demonstrated broad spectrum anticonvulsant activity, potentially through high receptor occupancy at $\alpha 2$ subunits

Potential Indications for Darigabat Beyond Epilepsy

Pipeline in a Pill

Potential for benzo-like activity with targeted GABA α 2/3/5 receptor selectivity

Benzos (Non-selective GABA Modulators)
Widely Prescribed for Seizures, Anxiety,
and Other Indications

Potential Large Indications Worldwide

▶	Epilepsy	~65M Patients
▶	Anxiety Disorders	~13M Patients (G7)
▷	Agitation	15-20M Patients
▷	Bipolar Disorder	~46M Patients



Significant need for GABA modulators that are better tolerated, less sedating, less addictive and supportive of chronic use

Darigabat TPP: Benzo-like Activity for Chronic Treatment

Darigabat Summary

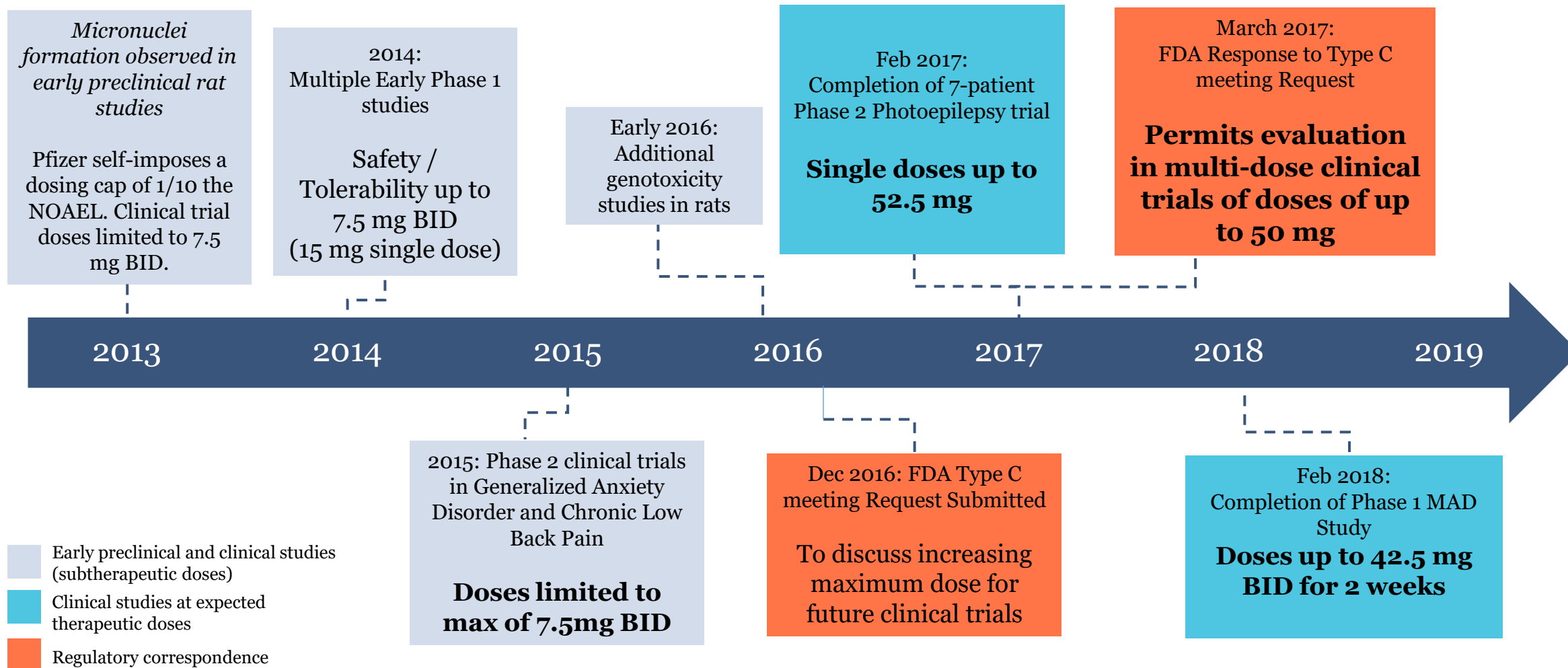
- 📈 Large markets (Focal & Generalized)
- 🔬 Novel mechanism
- ✓ Potential for better activity than chronic treatment alternatives
- ✓ Potentially favorable side effect profile
- 💰 Attractive pricing analogs

Pricing & Launch

- High branded sales despite many generics
- Branded US price analogs >\$10K/year
- Complex to change treatment in epilepsy
- 7-year+ average uptake in the category

History of Darigabat Development

- Results of early clinical trials were believed to be limited by Pfizer's self-imposed dosing cap



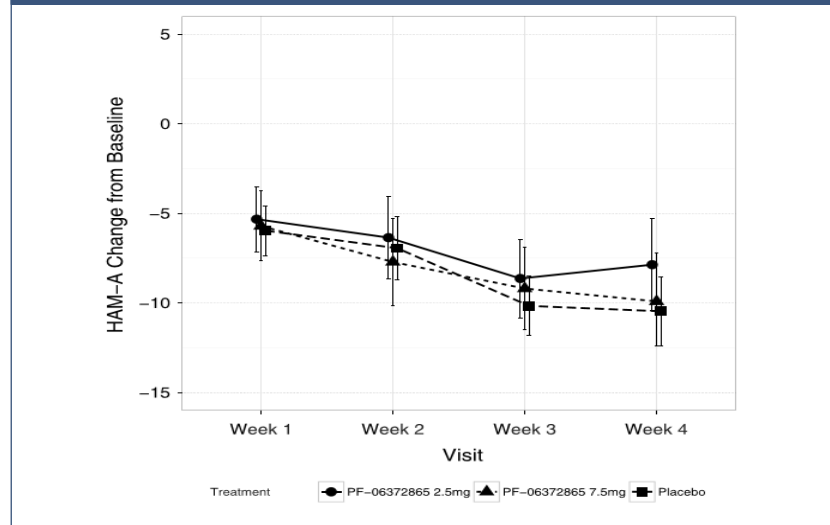
Prior Clinical Studies in Anxiety and Chronic Low Back Pain

Use of subtherapeutic doses and small sample size believed to account for lack of activity in prior trials

Phase 2: Generalized Anxiety Disorder

- Included drug-resistant patients on continued background treatment
- Sequential parallel comparison design
- Primary endpoint: HAM-A (minimum score at entry 20-22)
- 4 weeks on treatment: 2.5 mg BID darigabat, 7.5 mg BID darigabat, placebo
- Study stopped early for project prioritization - 90 enrolled of planned 384

Darigabat not differentiated from placebo on HAM-A



> 50% receptor occupancy remains unexplored in anxiety

Phase 2: Chronic Low Back Pain

- Included patients with > 6 months low back pain without radiculopathy
- Primary endpoint: change from baseline in low back pain score (measured on numerical rating scale)
- 4 weeks on parallel treatment: 7.5 mg BID darigabat, placebo, 500 mg naproxen BID (positive control)
- Study stopped early for futility following a planned interim analysis after 50% subjects had completed treatment

Darigabat not differentiated from placebo, naproxen



> 50% receptor occupancy remains unexplored in pain

Darigabat Favorable Side Effect & Tolerability Profile Across Trials

Darigabat has been tested in 289 subjects and was generally well-tolerated. There have been no clinically significant side effect observations from physical examination, vital sign measurements, laboratory safety assessments, or ECG parameters and no reports of sedation across single and multiple dose trials

I. Across Phase 1 trials:

- 81 healthy subjects received single doses of darigabat (0.04 to 100 mg); 55 healthy subjects received multiple doses of darigabat (2.5 to 42.5 mg BID)
- Most common AEs: dizziness, somnolence, and fatigue. All AEs across trials have been mild or moderate in severity
- No drug-related SAEs in Phase 1 trials
- Titration in multiple dose healthy volunteer studies appeared to reduce the incidence of somnolence and dizziness

II. Across Phase 2 trials:

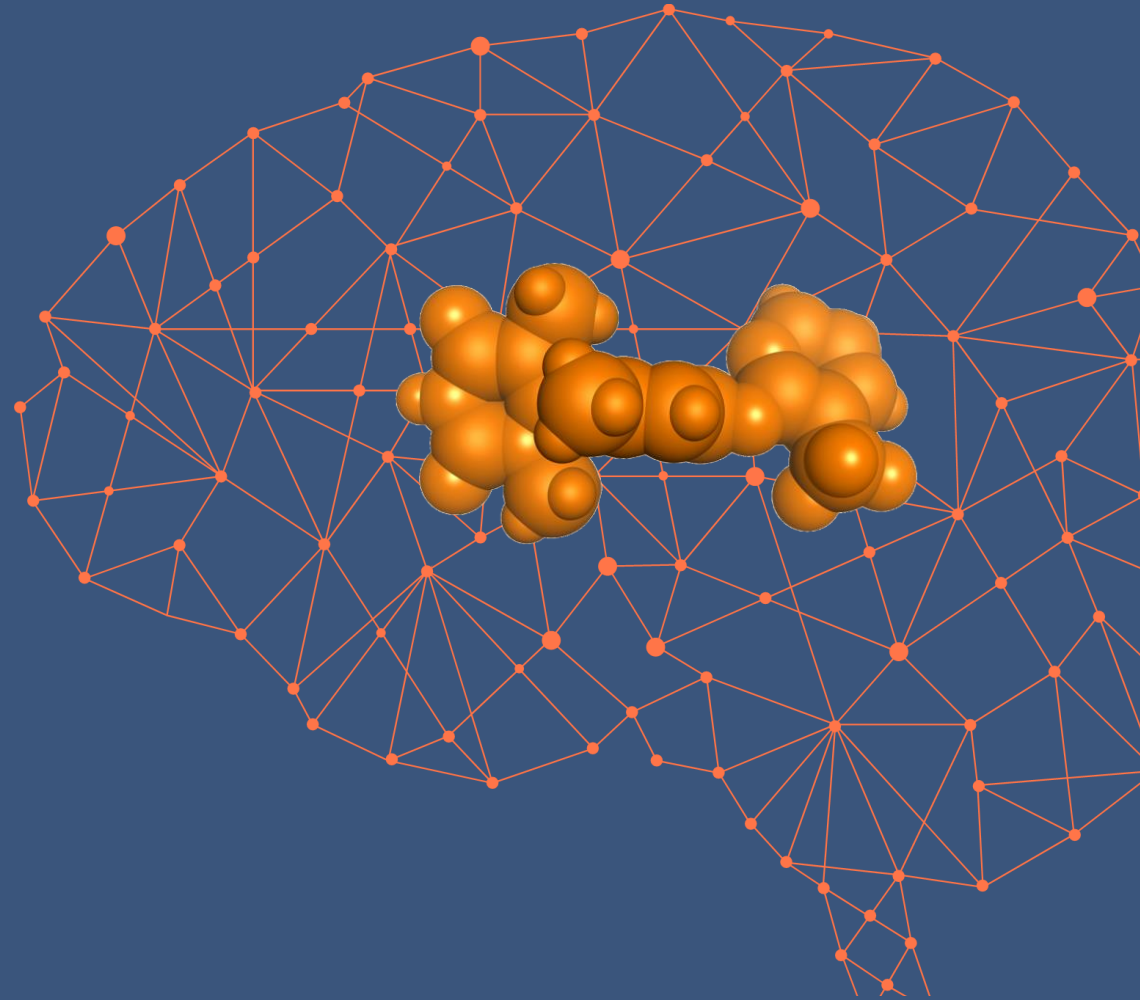
- 146 subjects received multiple doses of darigabat (2.5 to 7.5 mg BID); 7 subjects with documented photosensitive epilepsy received single doses of 17.5 mg and 52.5 mg in a crossover trial
- Most common AEs: dizziness and somnolence; the majority of AEs were mild or moderate
- In Study B7431007, there was limited increase in sleepiness as measured by the Epworth Sleepiness Score with either darigabat 7.5 mg, darigabat 2.5 mg or placebo at Week 2 and Week 4
- In Study B7431006, one patient experienced an SAE (transient ischemic attack) that was considered related to darigabat by the investigator. The patient had a history of high cholesterol levels and high blood pressure and was diagnosed with diabetes mellitus after the onset of TIA
- Use of titration in multi-dose Phase 2 trials appeared to mitigate CNS effects, including somnolence, over time

III. Other considerations:

- No evidence to date of withdrawal effects
- No evidence of the bone marrow effects seen in preclinical studies
- Reproductive effects are being addressed for all trials with requirements for contraception and standard warnings

Tavapadon in Parkinson's Disease

Additional Slides

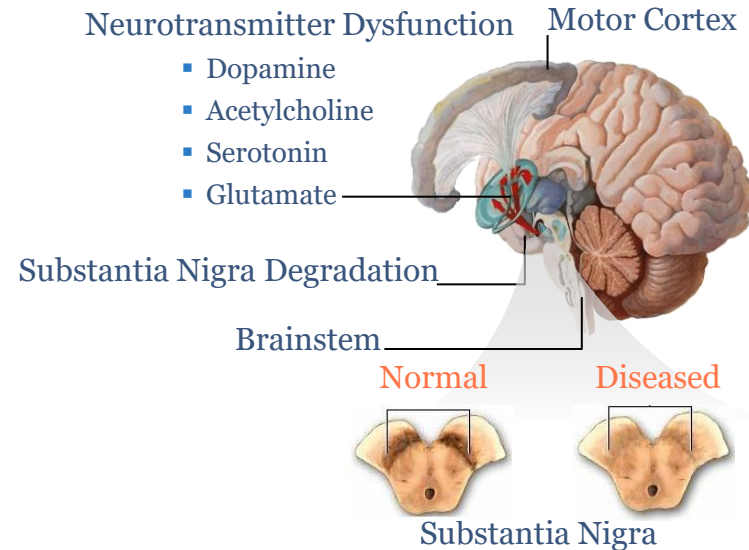


Parkinson's Disease Overview

Parkinson's disease is a progressive neurodegenerative disorder affecting regions of the brain that control balance and movement

Description

- Parkinson's disease is a degenerative neurological disorder characterized by progressive depletion of dopaminergic neurons in the substantia nigra region of the brain
- The lack of dopamine causes neurons to fire without normal control, leaving patients unable to control or direct their movement



Common Symptoms

- Symptoms of Parkinson's disease can be segmented into two categories – motor and non-motor:
 - Motor symptoms include tremor, decreased bodily movement (hypokinesia), slowness of movement (bradykinesia), stiffness and poor balance
 - Non-motor symptoms include cognitive dysfunction, psychosis, mood disorders, fatigue, *etc.*

Progression

- As symptom severity increases, patients often require increased doses of medication with decreasing efficiency, leading to “off” episodes
 - “Off” episodes are characterized by decreased motor function when patient's plasma drug levels fall below therapeutic levels
- Long-term levodopa use also leads to the development of dyskinesia or uncontrolled movement in PD patients

Genetic Indications

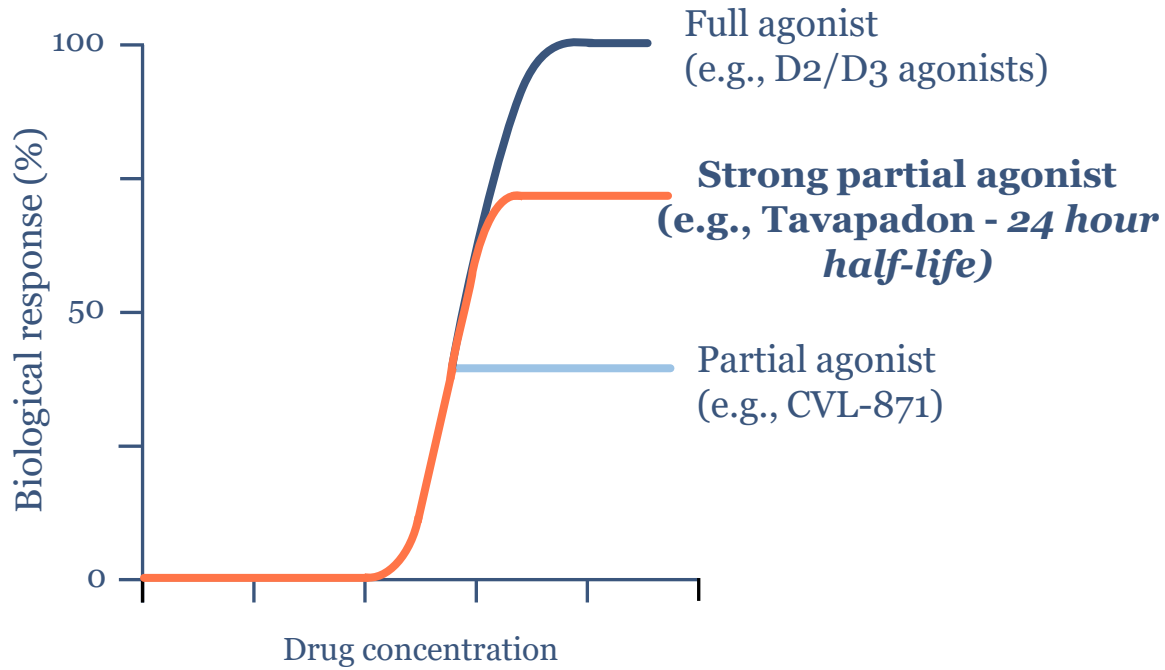
- Approximately 15% of Parkinson's patients have a family history of the disease. Such familial cases of the disease can be caused by mutations in the LRRK2, SNCA, PARK7, PINK1 or PRKN genes
- LRRK2 mutations attract greater attention from researchers since there are more known populations with this risk factor
 - G2019S is the most common LRRK2 mutation accounting for 3-6% of familial PD, and 1-2% of sporadic cases worldwide
 - This mutation is especially frequent in the Ashkenazi Jew and ArabBerber populations

Selectively Targeting Partial Agonism Designed to Improve Motor Control and Tolerability

D1/D5 Receptor Selectivity

D2/D3 Activation (Indirect Pathway)	Potential Effect	D1/D5 Activation (Direct Pathway)
+	Motor Control	++
	Cognition	++
	Motivation / Drive	++
-	Dose-Limiting Hypotension	
	Impulse Control Disorders	
	Sudden Daytime Sleepiness	

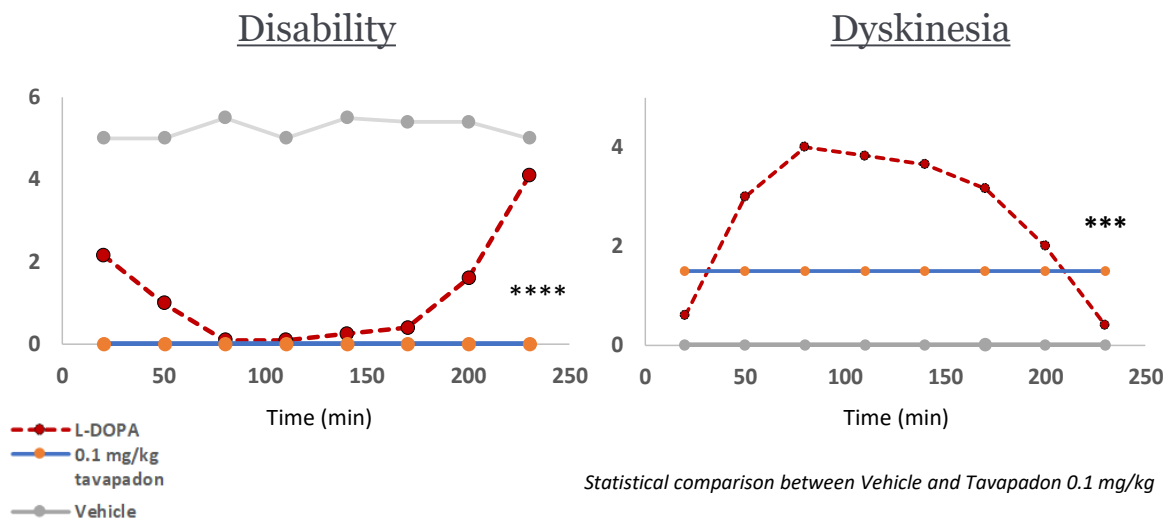
Degrees of Agonism (Illustrative)



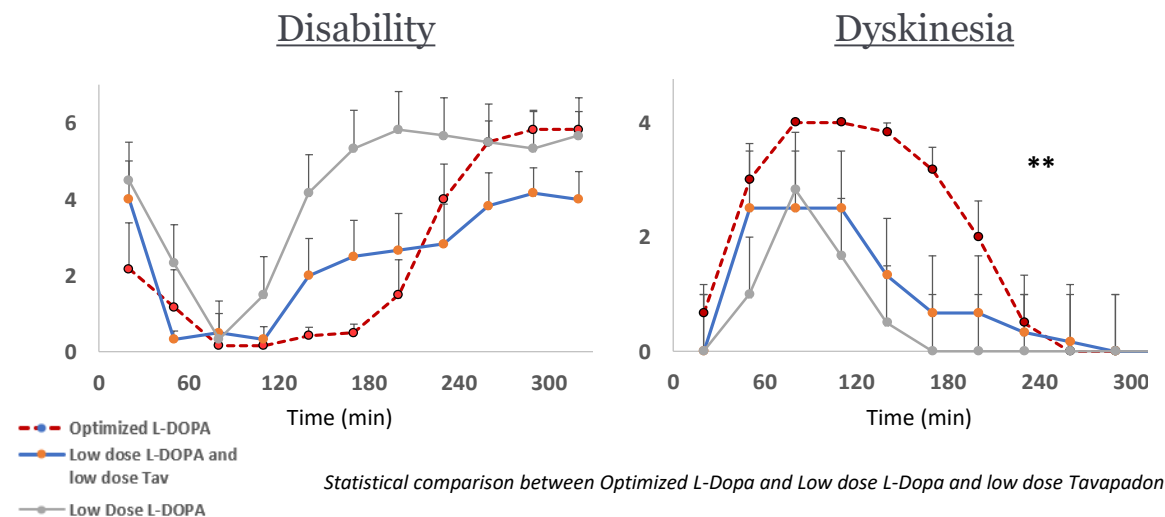
First Partial Agonist for Parkinson's → Avoids Dyskinesias

Unique and attractive combination of activity and low dyskinesias in highly translatable model of Parkinson's

Effect of Tavapadon vs. L-dopa on Disability and Dyskinesia



Potential to Reduce L-dopa Dose to Achieve Motor Control with Reduced Dyskinesia



Compared to L-dopa, tavapadon robustly reduced parkinsonian symptoms in the MPTP-lesioned primate model with a more **lasting effect** and **lower dyskinesia** levels

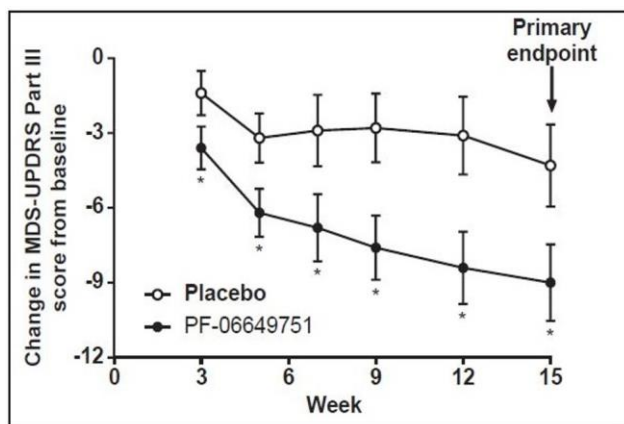
The combination of 33% L-dopa dose with 40% tavapadon dose showed **similar activity to L-dopa alone with statistically significant reduction in dyskinesia**

Phase 3 Program Designed to Show Superior Treatment Profile

Phase 2 Results Inform Phase 3 Design

Phase 2 Results

MDS-UPDRS III



MDS-UPDRS II+III

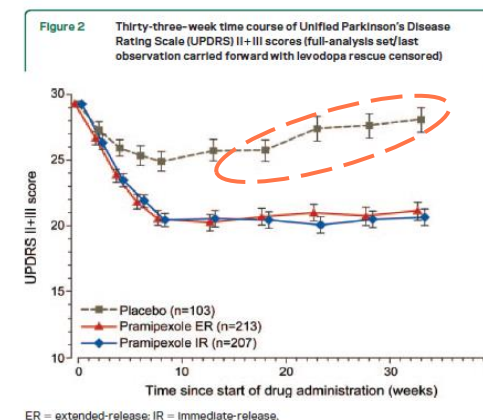
5.8 point improvement vs. placebo at week 15 on MDS-UPDRS II + III

Baseline score of 2 or greater on MDS-UPDRS II

~2 point improvement vs. placebo on Part II, excluding participants with baseline score of 0 or 1 in Phase 2*

Phase 3 Design

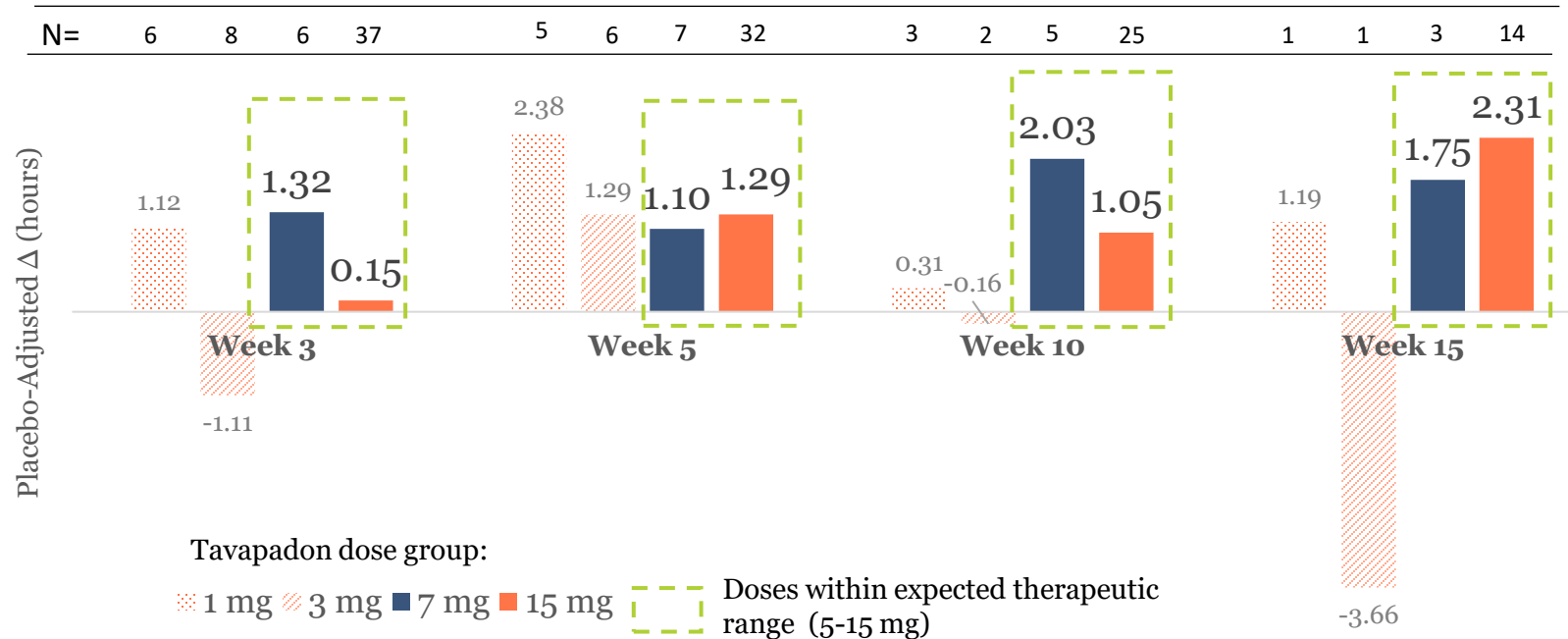
Placebo Attenuation at 6 Months



Further upside from forced titration to achieve maximum tolerated dose of tavapadon (less than 40% of patients were on the top dose in Phase 2)

Tavapadon Phase 2 Results: Clinically Meaningful ON-Time without Troublesome Dyskinesias

Tavapadon generally demonstrated at least 1 hour of improvement in ON-time without troublesome dyskinesias in late-stage PD



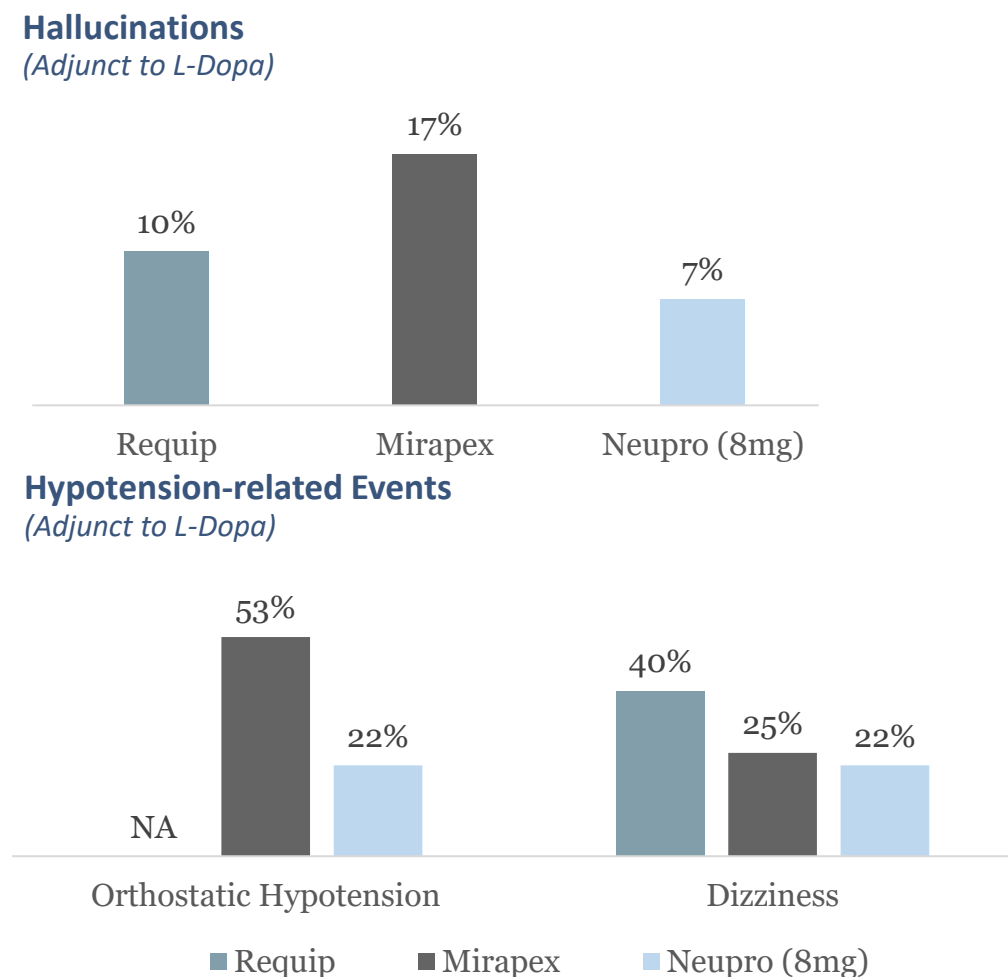
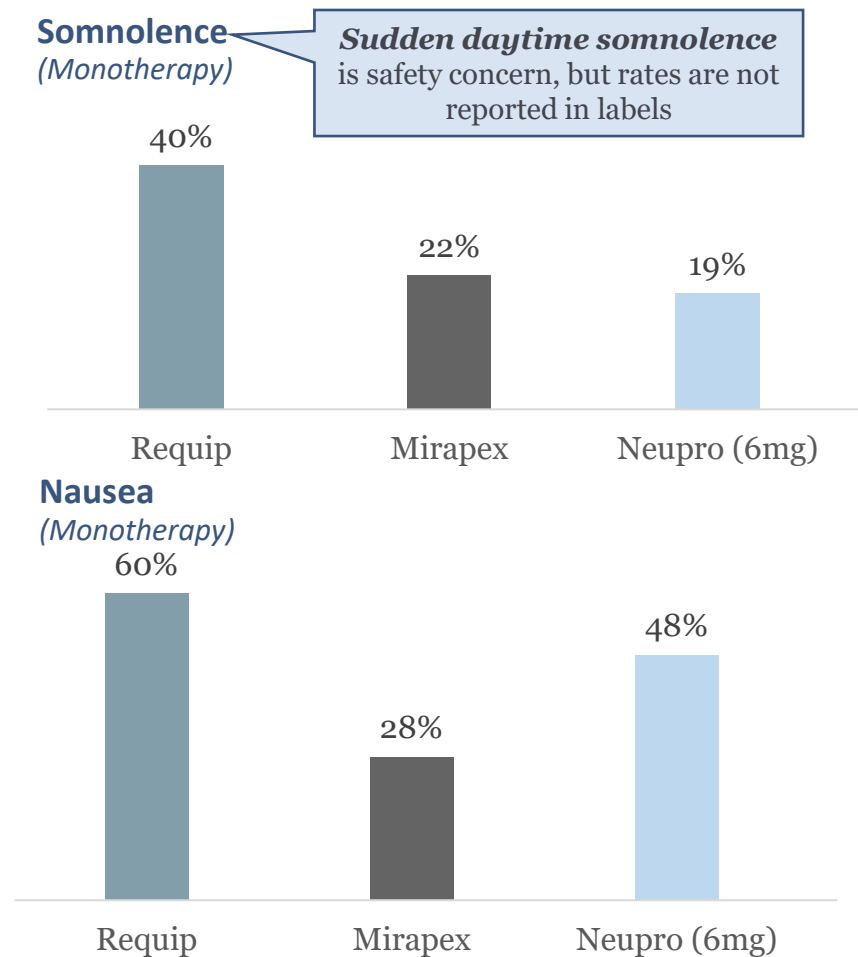
Note: Tavapadon showed -0.6 hour reduction on the primary endpoint, change in OFF-time at week 10



Tavapadon showed clinically meaningful benefit within therapeutic dose range of 5 mg to 15 mg

Historical D2/D3 Labels Show Significant Side Effect Profile

D2/D3 Side Effects: Sudden daytime somnolence, hallucinations, nausea and acute orthostasis





Overview of Tavapadon Clinical Trials To Date



Protocol ID	Phase	Trial End Date	N= (active/total)	Design
B7601001	Phase 1	7 Feb 2014	18/18	Single ascending dose (0.25-2.5 mg) in healthy volunteers (HV)
B7601002	Phase 1	16 Apr 2015	61/77	Multiple ascending dose study in HV (0.5-5 mg QD)
B7601007	Phase 1	04 Dec 2014	9/9	Single ascending dose (0.25 and 0.75 mg) with an antiemetic
B7601006	Phase 1	14 Sept 2017	11/11	CYP3A Victim DDI
B7601005	Phase 1b	10 Mar 2016	45/50	Open label multiple ascending dose (5/15/25 mg) in PD patients Adjunct with lowering of levodopa dose
B7601009	Phase 1b	28 Feb 2016	18/18	Placebo controlled single ascending dose (0.75/1/3/6/9 mg) in PD patients Monotherapy
B7601003	Phase 2	10 Nov 2017	85/108	Adjunct with levodopa (1/3/7/15 mg) in advanced PD patients (w/ OFF-time \geq 2.5h at baseline) Three week dose titration, 15 weeks total dosing
B7601011	Phase 2	29 Jan 2018	29/57	Monotherapy in newly diagnosed PD patients; flexible dosing Seven week dose titration, 15 weeks total dosing

Thank you

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