



Innovative Treatments for Central Nervous System Disorders

February 2021

Allosteric modulators for human health

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Addex Overview

3 clinical programs	<ul style="list-style-type: none">• Dipraglurant PD-L1D study 301 starting H1 2021*• Dipraglurant blepharospasm Phase 2 study starting H1 2021• ADX71449 (J&J) epilepsy Phase 2 study starting Q2 2021
Leading allosteric modulator technology platform	<ul style="list-style-type: none">• Allosteric modulation is a validated & differentiated pharmacological approach to address drug targets• Proprietary biological screening assays and chemical library
In house discovered pipeline	<ul style="list-style-type: none">• Multiple novel drug candidates entering clinical candidate selection• Driving long term growth & future partnership opportunities
Partnerships	<ul style="list-style-type: none">• J&J - €109M in milestones & double digit royalties• Indivior - \$330M in milestones, royalties up to double digit & funded research program
Top tier US investors	<ul style="list-style-type: none">• Dual listed on SIX Swiss Exchange & US Nasdaq Capital Market• Cash of CHF18.7M at 31 Dec 2021• Completed \$11.5M capital increase 11 Jan 2021

Addex Pipeline

Molecule / MoA	Indication	Partner	Pre-clinical	Phase 1	Phase 2	Phase 3	Milestone
Dipraglurant (mGlu5 NAM)	PD-LID						Data Q4 2022
	Blepharospasm						Data Q4 2021
ADX71149 (mGlu2 PAM)	Epilepsy						Data H1 2022
GABA _B PAM	Addiction						
	CMT1A						
mGlu7 NAM	PTSD						
mGlu2 NAM	Mild neurocognitive disorders						
mGlu4 PAM	Parkinson's disease						
mGlu3 PAM	Neurodegenerative disorders						

Lead Program Entering US Pivotal Study

Experienced Team

Leadership Team

<p>Tim Dyer CEO / CFO</p> <p>Co-Founder of Addex Formerly with PwC UK Chartered Accountant</p>	<p>Dr Roger Mills Chief Medical Officer</p> <p>Developed Nuplazid in PD Psychosis 30 years in Pharma industry including Pfizer, Gilead and Acadia</p>	<p>Dr Robert Lutjens Head of Discovery Biology</p> <p>Member of Addex founding team Formerly with Glaxo & Scripps Research Institute</p>	<p>Dr Jean-Philippe Rocher Head of Discovery Chemistry</p> <p>Member of Addex founding team Formerly with Pierre Fabre, GSK and Mitsubishi</p>
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Non-executive Directors

<p>Vincent Lawton Chairman</p> <p>Former European Head of Merck & Co. Former MHRA Board member</p>	<p>Ray Hill Board member</p> <p>Former Executive Director Merck & Co.</p>	<p>Jake Nunn Board member</p> <p>Venture advisor and former Partner at New Enterprise Associates</p>	<p>Isaac Manke Board member</p> <p>General Partner at Acorn Bioventures. Formerly Partner at New Leaf Venture Partners</p>
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Scientific Advisory Board

<p>Darryle Schoepp Chairman of SAB</p> <p>Former leader of Neuroscience research department at Eli Lilly, and at Merck was Neuroscience research therapeutic area leader</p>	<p>Mark Bear Picower Prof. of Neuroscience at MIT</p> <p>Formerly on faculty of Brown University School of Medicine and an Investigator of the Howard Hughes Medical Institute</p>	<p>Peter Bernstein Principal, PhaRmaB LLC</p> <p>Formerly with ICI Astra Zeneca Awarded numerous accolades including Fellow of the American Chemical Society</p>	<p>Benny Bettler Biomedicine Prof. at Basel University</p> <p>Formerly at Novartis and discovered allosteric modulators at GABA_B receptor and recipient of the Peter Speiser Award</p>
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Milestones

Milestone	Timing
Dipraglurant for PDLID	
Phase 2b/3 - start study	H1 2021
Phase 2b/3 - topline results	Q4 2022
Dipraglurant for Blepharospasm	
Phase 2a - start study	H1 2021
Phase 2a - topline results	Q4 2021
ADX71149 for Epilepsy	
Phase 2a - start study	Q2 2021
Phase 2a - topline results	H1 2022
GABA _B PAM for Addiction and CMT1a	
Complete clinical candidate selection	Q4 2021
Start IND enabling studies	Q1 2022

Dipraglurant for Levodopa-Induced Dyskinesia in Parkinson's Disease (PD-LID)

Compelling Rationale to Develop Dipraglurant for PD-LID

- Large underserved patient population in need of improved treatment options
- Significant commercial opportunity with limited competition
 - 1M Parkinson's disease patients in US of which >170,000 have dyskinesia
 - GOCOVRI® price: \$34K p.a., Nuplazid® price: \$45K p.a.
 - US LID market estimated at \$4B
- Strong mechanistic rationale for blocking mGlu5 to inhibit glutamate signalling
- Supportive pre-clinical data and Phase 2 clinical data
- PK profile ideally suited for treatment of LID
- Dipraglurant is active on same biological pathway as amantadine (inc. GOCOVRI®)
 - Decreases glutamatergic tone
 - Unlike amantadine, dipraglurant:
 - Restores synaptic plasticity to prune aberrant signalling
 - Highly selective with limited off target activity
- Novartis mGlu5 NAM (mavoglurant) data supportive of mGlu5 target & rationale for dipraglurant PK profile

Disability and Impact of PD-LID

Invariably associated with long-term L-dopa use

- Dyskinesias caused by neurodegeneration
- Dopamine replacement lowers the triggering threshold for symptoms
- LID can become as disabling as the PD symptoms themselves

Symptoms include dystonia, chorea, and choreoathetosis

- Uncontrollable muscle contractions, twisting and writhing
- Painful and severely disabling
- Causes fatigue/exhaustion and increased risk for falls and injuries
- Social withdrawal, reduced quality of life and increased burden on caregiver

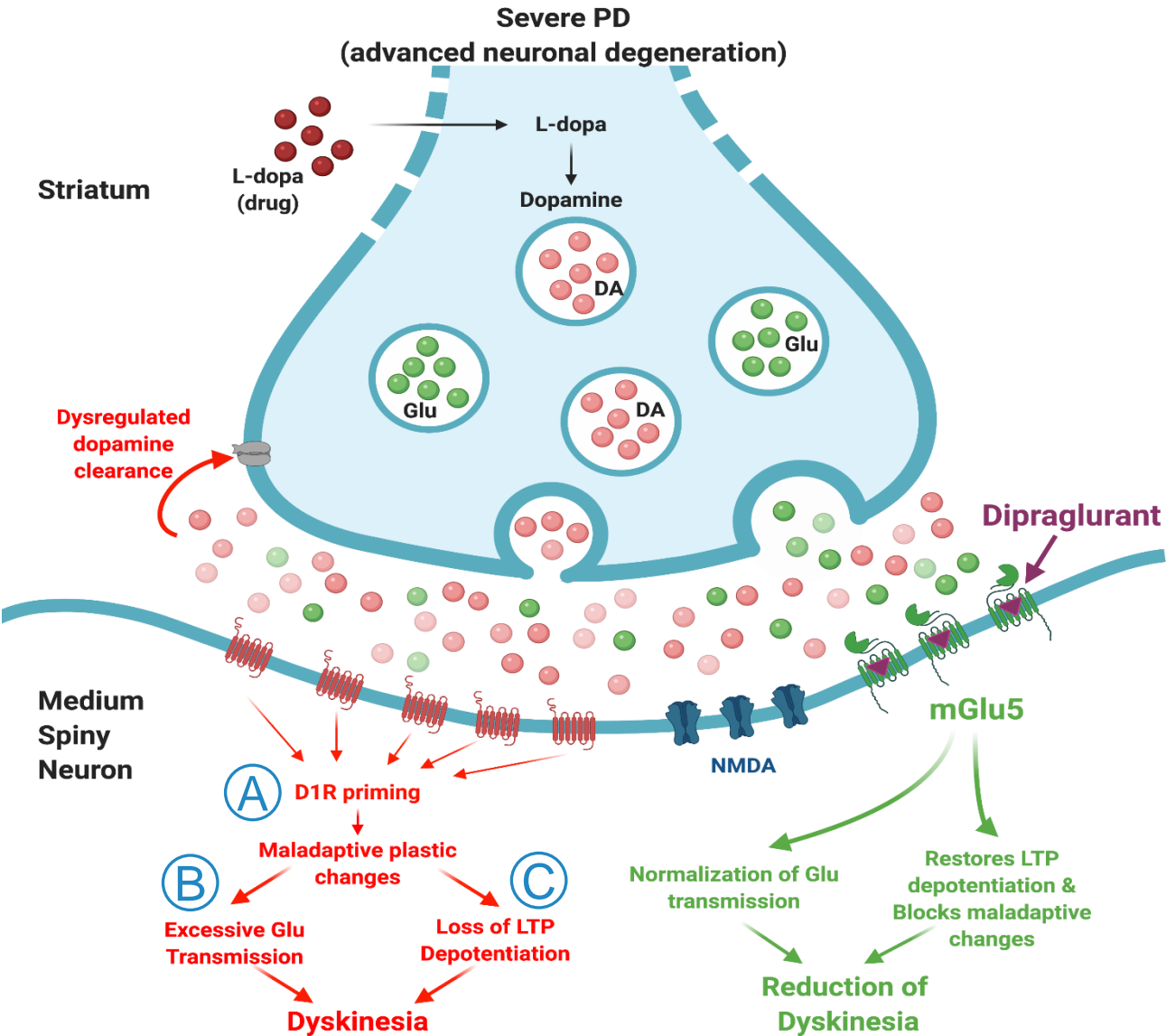
Prevalence related to disease duration

- >40% of patients experience LID within 4-6 years of L-dopa treatment
- Increases to 90% after 9 -15 years
- Patients treated with next-generation L-dopa will still experience LID

PD drug efficacy wanes over time - exacerbated by emergence of LID

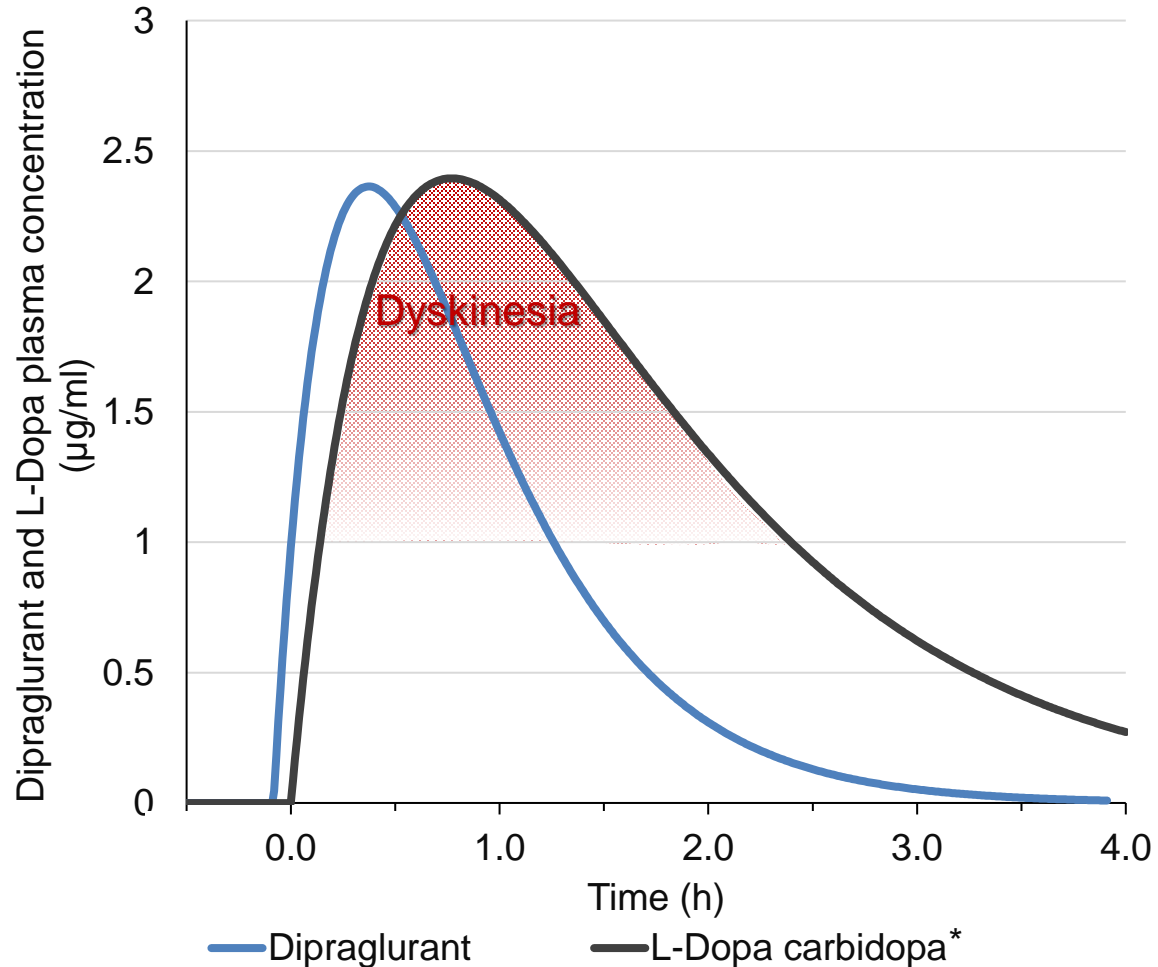
Treatment becomes a balancing act requiring constant adjustments to ensure symptom control & minimize intolerable side effects

MoA Rationale for Targeting mGlu5 Inhibition in PD-LID



- Loss of substantia nigra neurons combined with the non-physiological, pulsatile stimulation of dopamine receptors with L-dopa are at the basis of LID development
- In the striatum, LID is the result of:
 - A** *D1 receptor priming*
 - B** *Excess glutamate transmission*
 - C** *Loss of LTP depotentiation*
- mGlu5 receptor is an attractive target due to its modulatory action - normalizing glutamatergic activity and restoring LTP depotentiation
- Inhibiting mGlu5 decreases excess glutamatergic tone thereby controlling dyskinesia
- Dipraglurant is an oral, highly selective NAM of the mGlu5 receptor

Dipraglurant PK is a Key Advantage for Treating LID

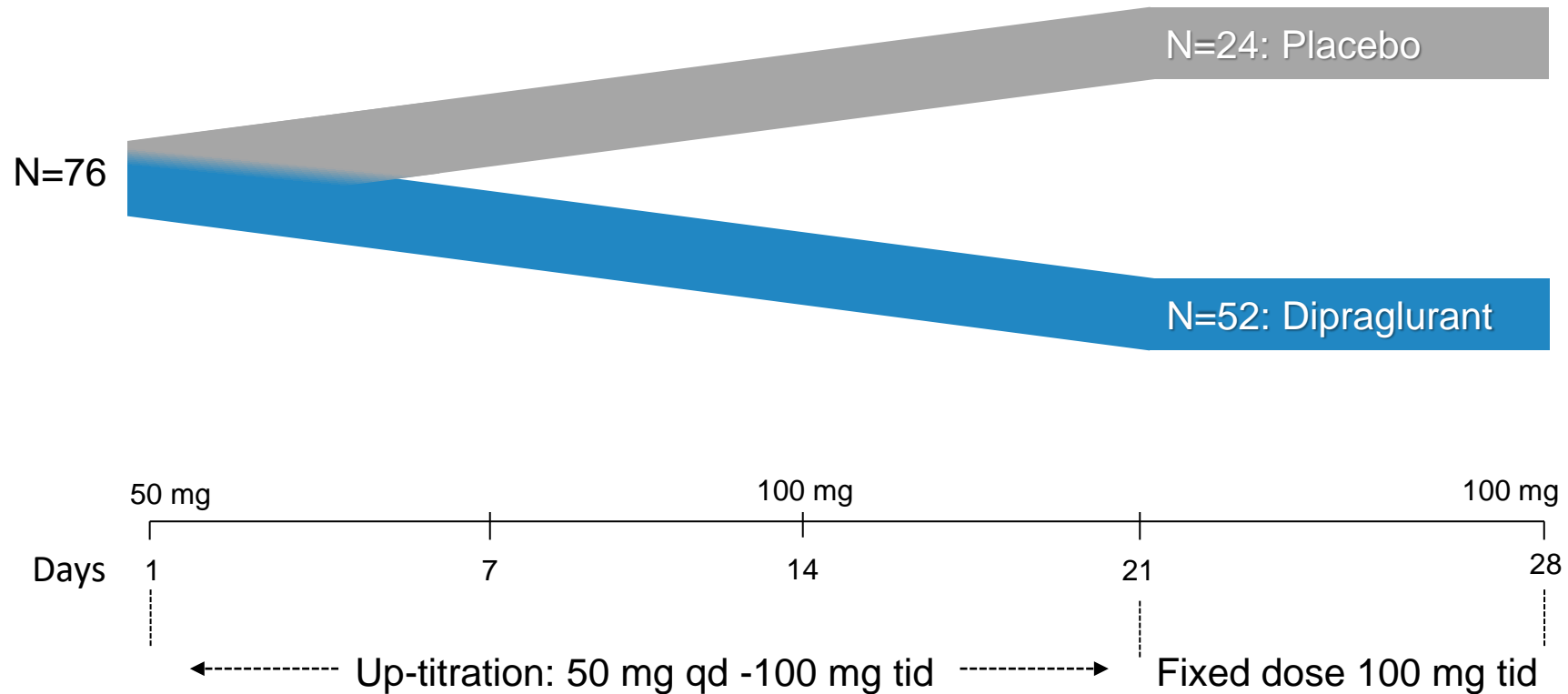


- Dyskinesia symptoms are correlated to peak levels of L-dopa
- PK of dipraglurant allows control of glutamatergic tone ahead of L-dopa Cmax

Dipraglurant normalizes abnormal glutamate stimulation during peak levodopa dose

Dipraglurant peaks ahead of L-dopa for optimal LID control

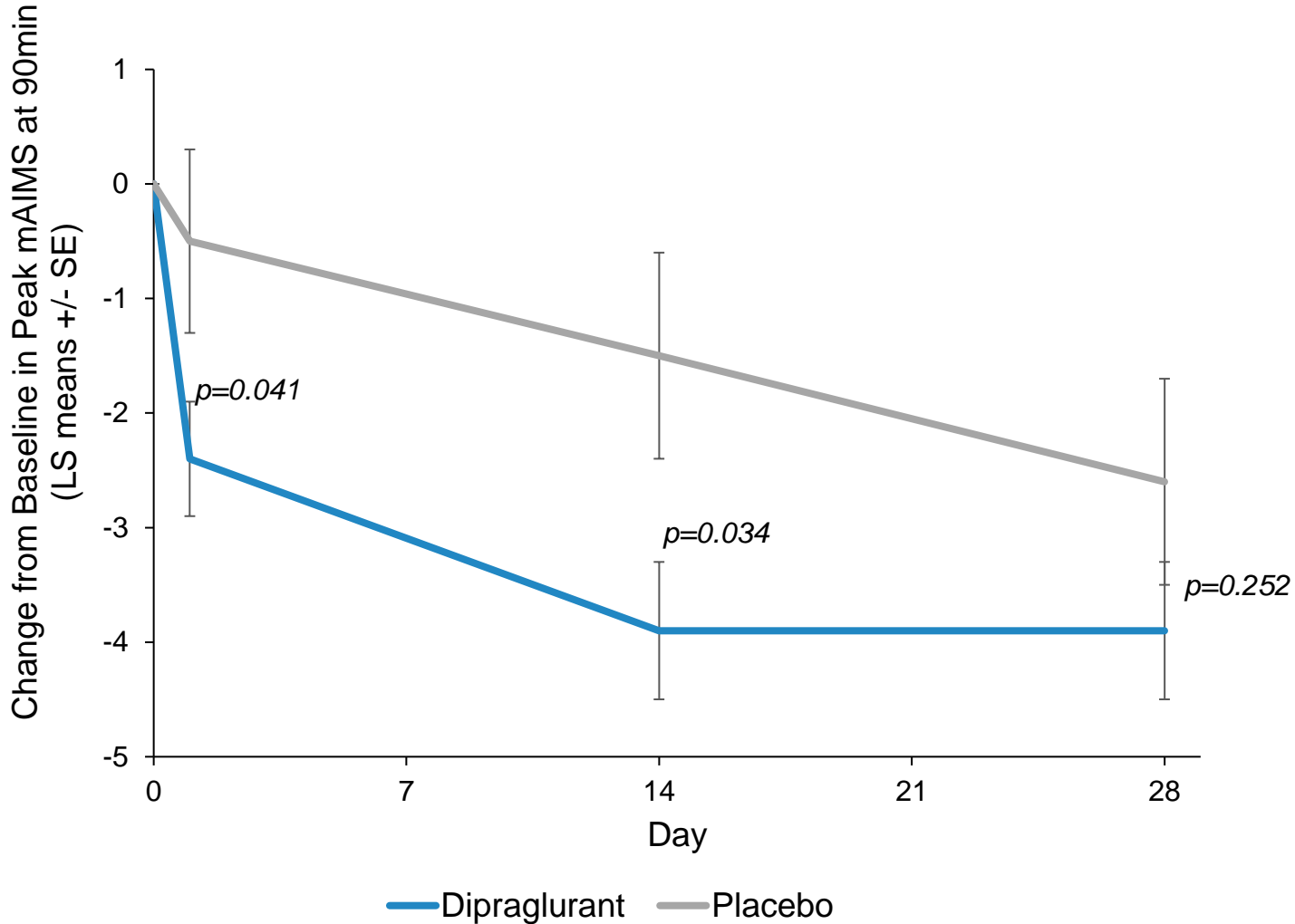
Dipraglurant Phase 2a Study in LID (in US and Europe)



- **Primary objective:** safety & tolerability
- **Secondary objective:** exploratory efficacy:
 - Modified Abnormal Involuntary Movement Scale (mAIMS) on days 1, 14 and 28
 - Clinician Global Impression of Change (CGIC)
 - Patient diaries of “On” & “Off” time

Measured acute effect of mid-day dose on days 1, 14 and 28

Dipraglurant Improves LID by 30%

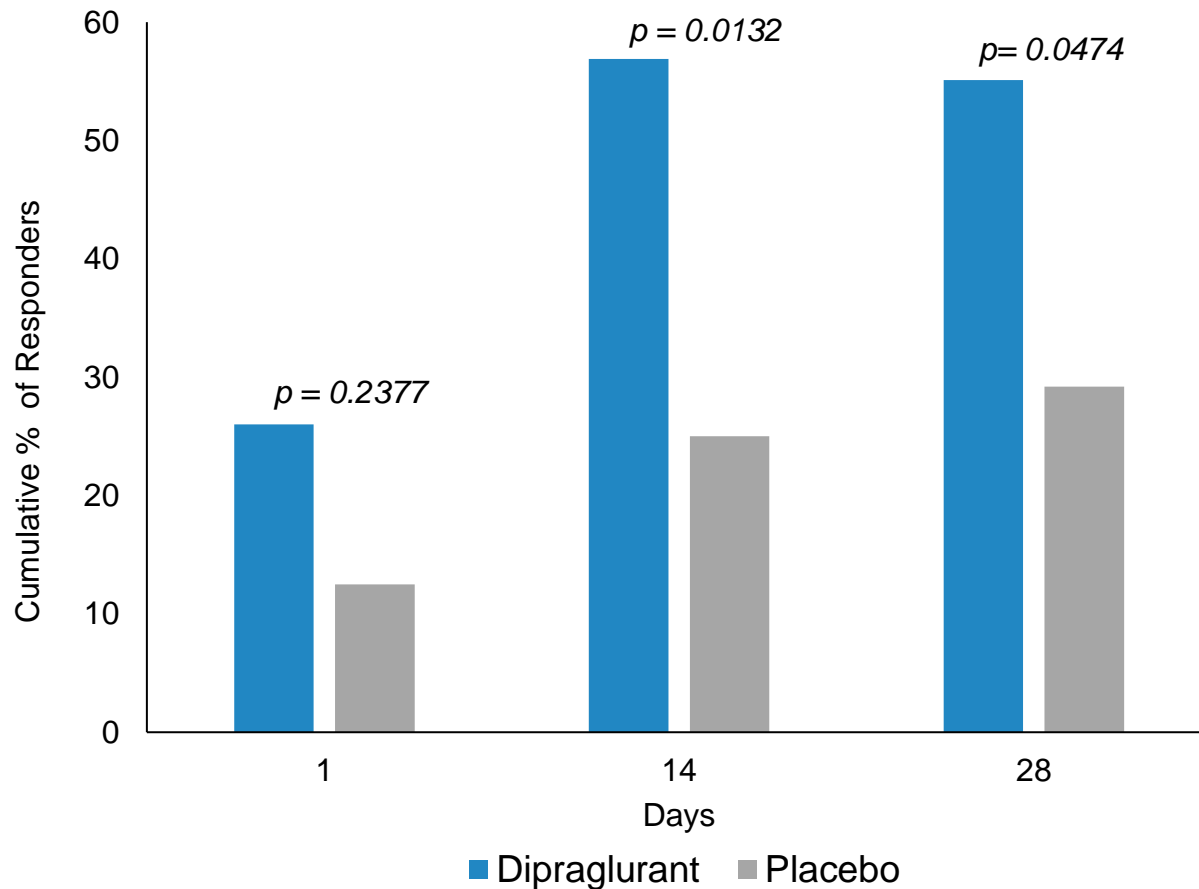


- Statistically significant effects: Day 1 (50mg) and Day 14 (100mg)
- Improvement maintained through Day 28
- Increasing placebo response caused significance to be lost at Day 28
- No placebo mitigation in study

Mean % change of peak mAIMS from baseline		
Midday dose	Dipraglurant	Placebo
Day 1 (50 mg)	19.9%	4.1%
Day 14 (100 mg)	32.3%	12.6%
Day 28 (100 mg)	31.4%	21.5%

Responder Analysis Demonstrates Dipraglurant Significant Benefit

Percent of patients with $\geq 30\%$ improvement on mAIMS

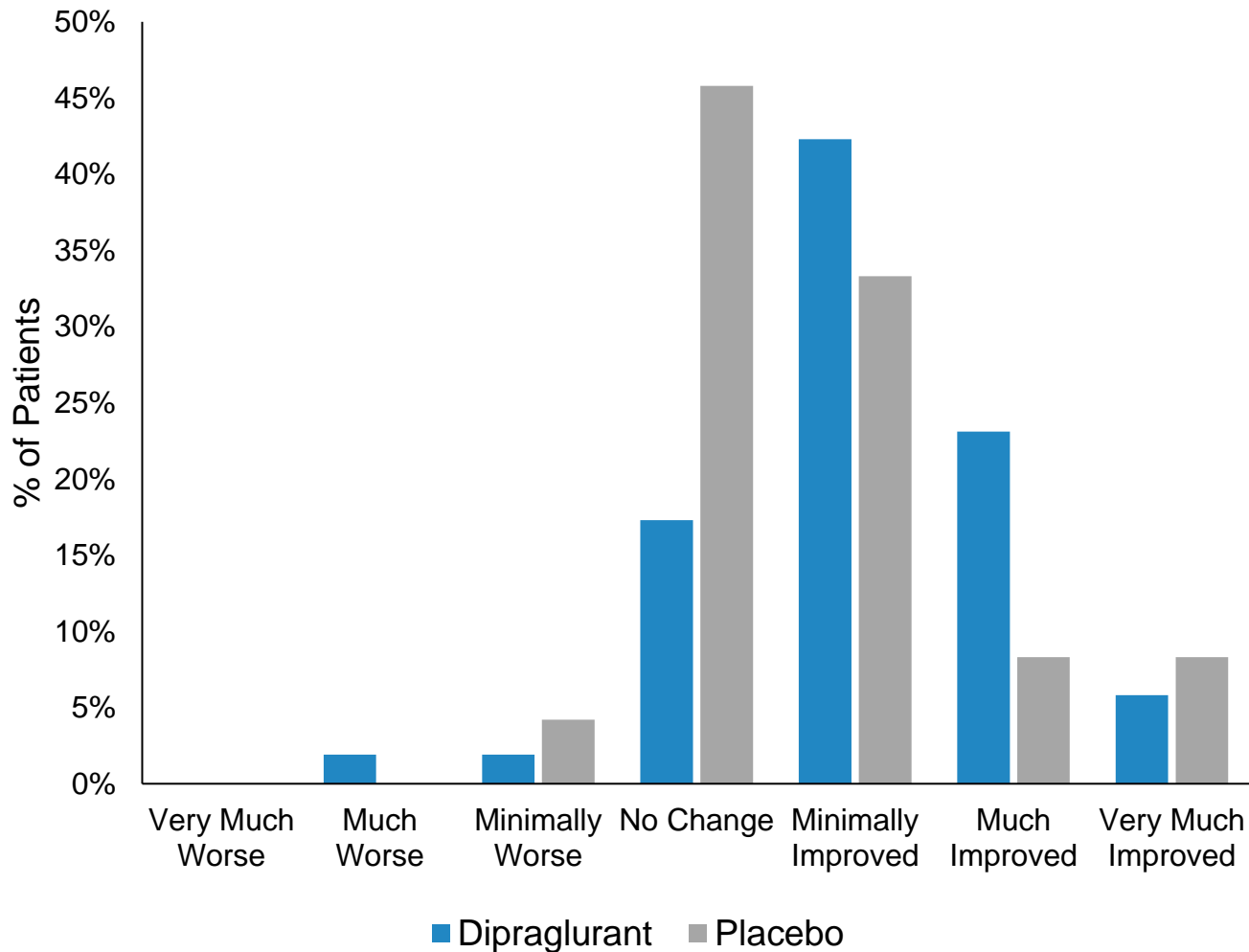


Responder analysis ($\geq 30\%$ change of mAIMS from baseline)				
Midday dose	Dipraglurant		Placebo	
Day 1 (50 mg)	n=13	26.0%	n=3	12.5%
Day 14 (100 mg)	n=29	56.9%*	n=6	25.0%
Day 28 (100 mg)	n=27	55.1%*	n=7	29.2%

*statistically significant

Reinforces robustness of dipraglurant anti-dyskinetic effect

Significant Improvement on CGI-C



	Dipraglurant	Placebo
Improved (p<0.05)	71.2%	49.9%
No change	17.3%	45.8%

- Simple scale reflecting clinical assessment by treating physician
- More objective than mAIMS
- Assessed at end of study compared to baseline
- Supports use of UDysRS in pivotal program

Dipraglurant Demonstrated Good Safety and Tolerability in PD Patients

- Adverse events common in both treatment groups (dipraglurant 88.5%, placebo 75%)
- Most common AEs:

	Dipraglurant	Placebo
Worsening Dyskinesia	21% (15.3%*)	12.5%
Dizziness	19%	12.5%
Nausea	19%	0%
Fatigue	15%	4%

* 3 of 11 AEs of “worsening dyskinesia” occurred in the follow up period (i.e., after drug discontinuation). On treatment incidence = 15.3% dipraglurant, 12.5% placebo

- AEs led to discontinuation in 2 patients (dipraglurant 100 mg)
- Fewer AEs at 50 mg (Weeks 1 and 2) 53% vs 58% placebo compared to 100 mg (Weeks 3 and 4) 73% vs 63% placebo
- No treatment effects on ECG, HR, BP, haematology and biochemistry

Safety profile supports continued development in PD-LID (KOLs and DSMB)

UDysRS: An Improved and Validated Dyskinesia Rating Scale

	UDysRS	mAIMS
Characteristics	<ul style="list-style-type: none">• Recommended by Movement Disorder Society (MDS)• FDA regulatory precedent (GOCOVRI® approval)• Contains anchored objective clinician evaluated measures of dyskinesia• Includes both patient and physician assessments of impairment• Less prone to placebo effect	<ul style="list-style-type: none">• Suboptimal for detecting treatment-related changes• Limited to patient assessments• Prone to placebo effect
Clinimetrics	<ul style="list-style-type: none">• Validated	<ul style="list-style-type: none">• Only the original version has been validated
Development	<ul style="list-style-type: none">• Developed in 2009 specifically for dyskinesia in PD	<ul style="list-style-type: none">• Developed in 1970 for tardive dyskinesia in psychiatry

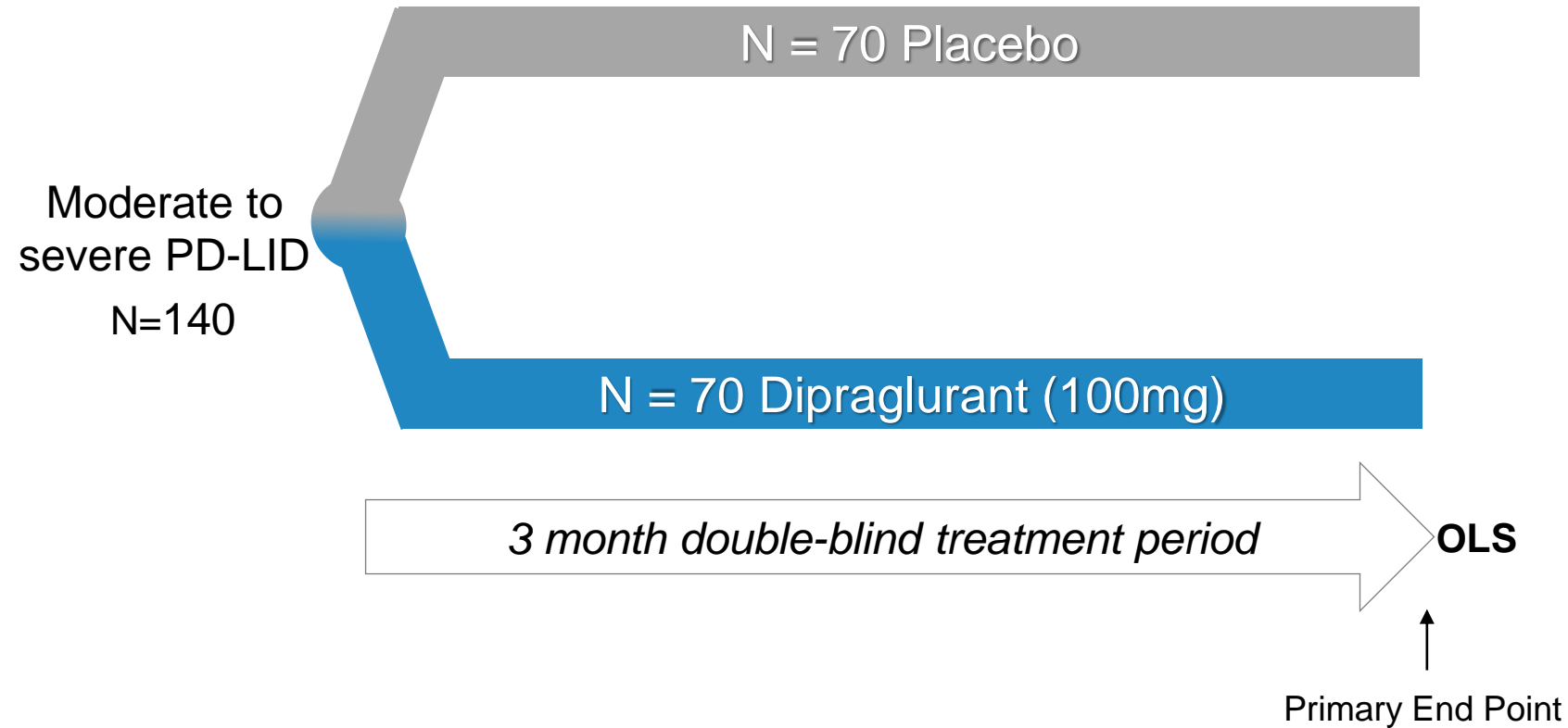
Dipraglurant PD-LID Studies – Management of Placebo Response

- Use of UDysRS
 - More sensitive to changes in LID
 - Less prone to placebo response
- Raters will be qualified by the MDS
 - Expert rater review to further ensure quality
- Requirement for moderate to severe symptom scores at screening and baseline
- BPST-Dys (non-pharmacologic intervention) to be used during screening
- Longer 12-week treatment period expected to mitigate placebo response

Dipraglurant PD-L1D Registration Program Started

- Pivotal registration program ongoing
- Study 301 expected to start H1 2021*
 - Data read-out expected Q4 2022
 - Primary endpoint: UDysRS
 - Placebo mitigation is a priority
- 12-month Open Label Study (302) starting in parallel to study 301
 - 6- and 12-month safety data
- Second pivotal registration study (303) to follow study 301 completion

Dipraglurant First Pivotal PD-LID Study (301)



- Primary objective: Efficacy in reducing LID
 - UDysRS change from baseline at 3 months
- Secondary objectives
 - CGI-S
 - MDS-UPDRS Part III change from baseline
 - Patient diaries, on & off time
 - Safety and tolerability

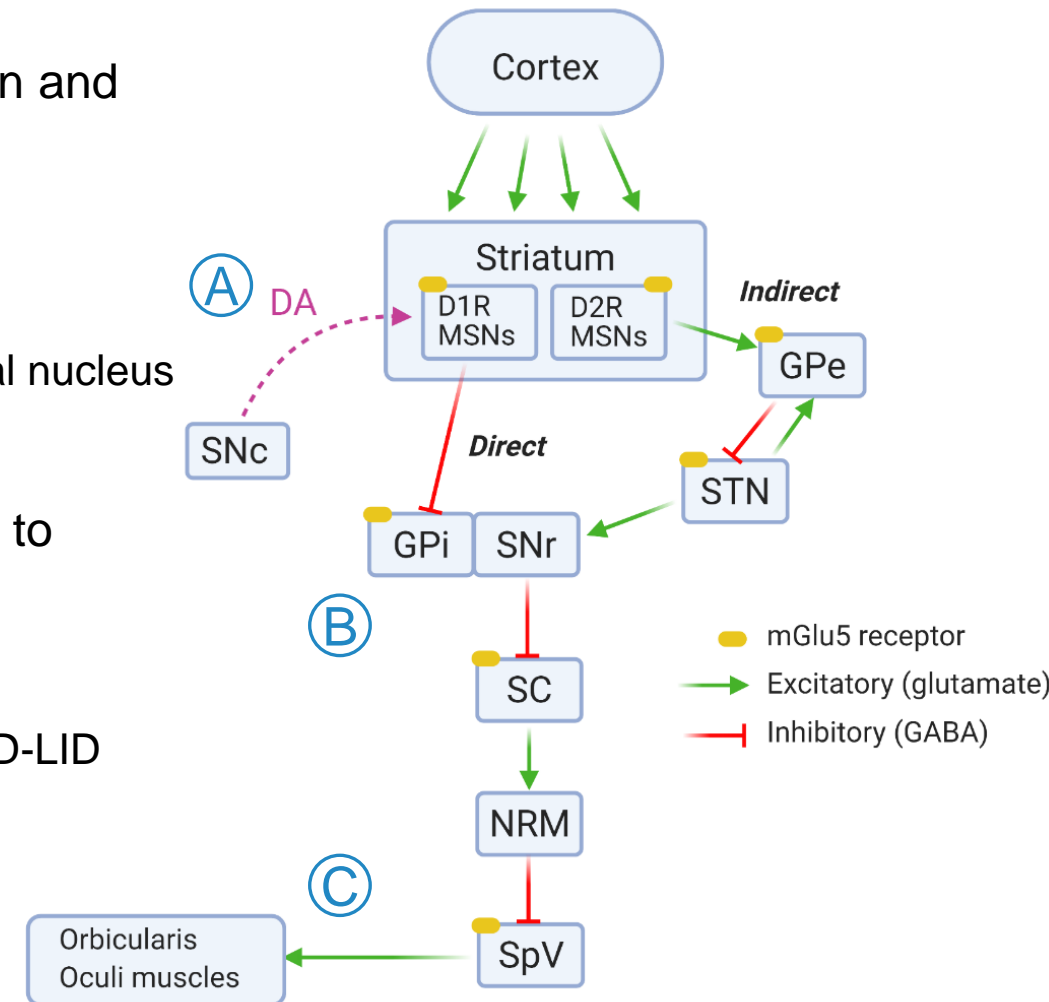
Dipraglurant for Dystonia – Blepharospasm

Blepharospasm (BSP)

- Type of dystonia affecting eyelid muscles
 - Results in sustained eyelid closure causing substantial visual disturbance or functional blindness
 - >50% of BSP patients symptoms spread to other cranio-facial muscles
- At least 50,000 BSP patients in US, ~2000 new patients diagnosed annually
- Botulinum toxin (BoNT) injections are the only approved treatment
- Surgical approaches including myectomy are invasive and frequently not of benefit
- Phase 2 feasibility study in BSP with dipraglurant IR expected to start in H1 2021 and read out data by the end of 2021
- Dipraglurant extended release (ER) formulation being developed
- Phase 2a proof of concept with dipraglurant ER planned for 2022
- Potential to expand to other dystonias

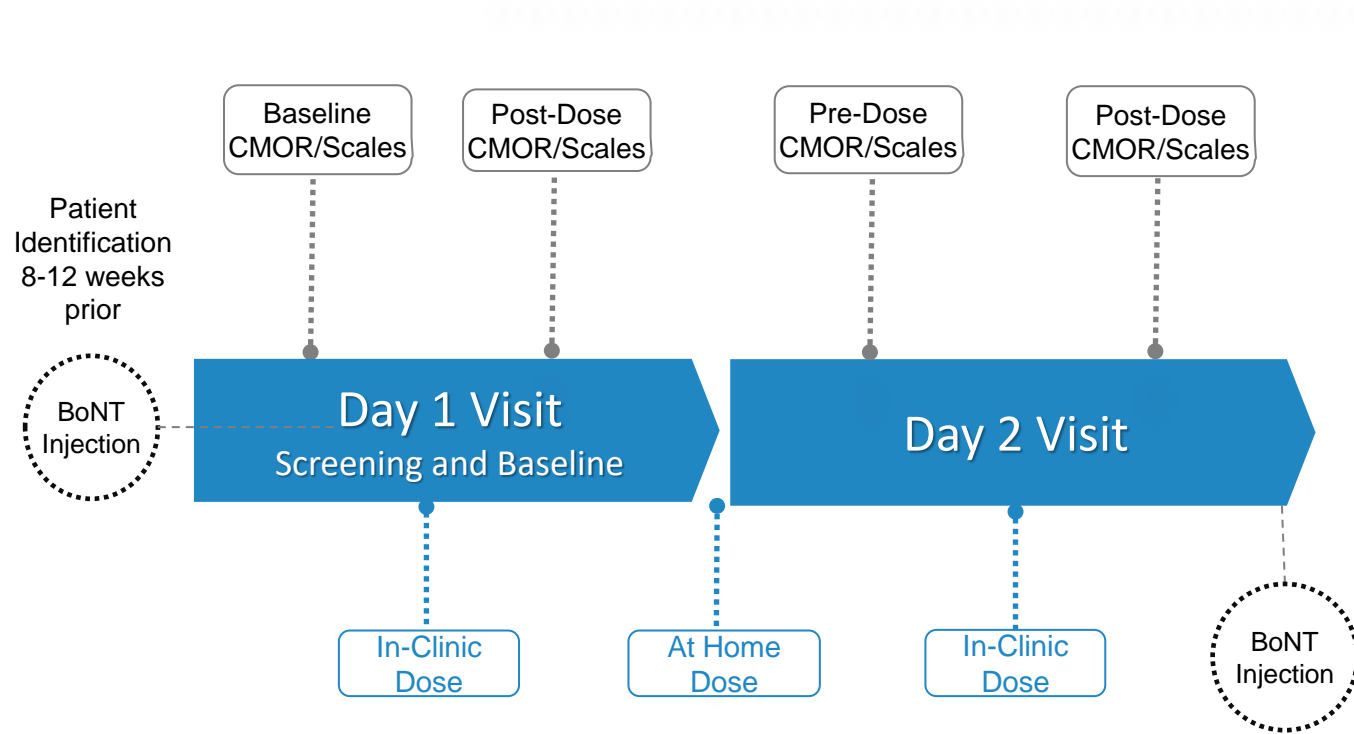
Rationale for Targeting mGlu5 Inhibition in Dystonia & BSP

- Dystonias are *neuro-functional* rather than *neuro-degenerative*
- Common features include alterations in neuronal connectivity/function and synaptic communication
- BSP pathophysiology is linked to:
 - Ⓐ Reduction of dopamine input into striatum
 - Ⓑ Increased inhibition of direct pathway from superior colliculus to trigeminal nucleus
 - Ⓒ Overexcitation of the signal leading to blink reflex
- Pathogenesis involves aberrant or maladaptive brain plasticity linked to excessive sensory stimulation and/or repetitive motor tasks
- Dipraglurant shows robust preclinical validation:
 - Dose-dependent reduction of dystonia in MPTP-lesioned NHP model of PD-LID
 - Effective in tottering mouse model of generalized dystonia
 - Reverses synaptic plasticity alterations observed in two distinct genetic models of dystonia (DYT1 mice & DYT25 rats)
- Dipraglurant has shown anti-dystonic effect in PD patients



Adapted from Peterson & Sjenowski, 2017

Blepharospasm Phase 2 Feasibility Study Design



- Patients with benign essential BSP, who experience moderate/severe symptoms prior to their regular dose of BoNT
- Single center, randomized, double-blind, placebo controlled
- Approx. 15 patients
- Dipraglurant IR - 50mg, 100 mg and placebo
- Efficacy endpoints include:
 - Computational Motor Objective Rater (CMOR)
 - Clinician rating scales
 - Patient reported outcomes

ADX71149 (JNJ-40411813) for Epilepsy Partnered with Janssen (JNJ)

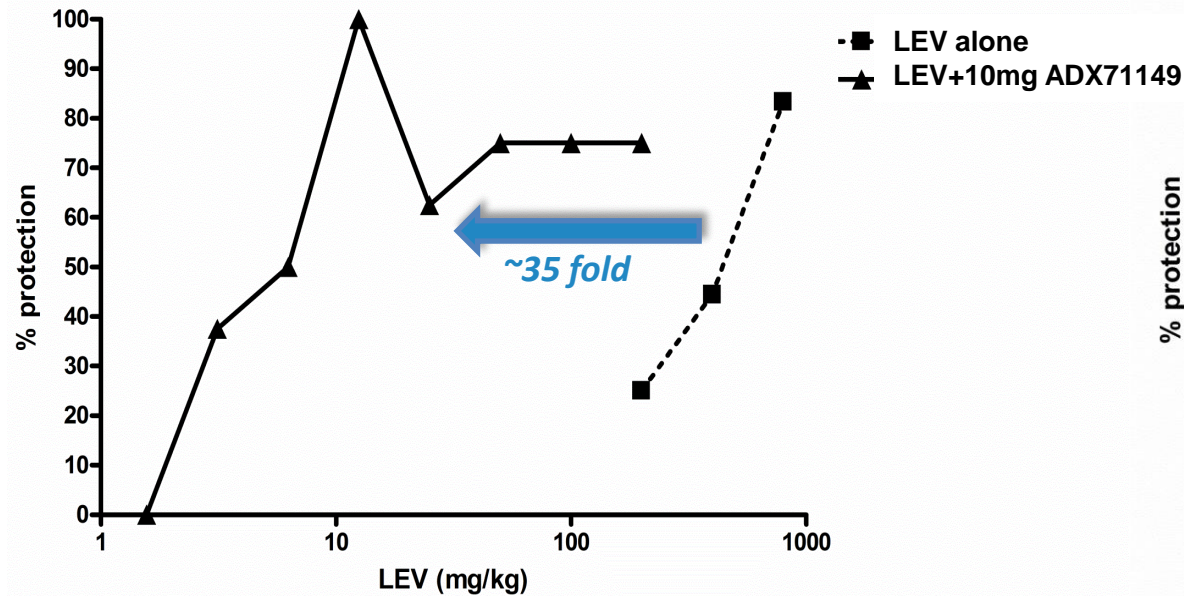
ADX71149 Opportunity in Epilepsy

<h2>Large market & unmet medical need</h2>	<ul style="list-style-type: none">• Market projected to reach \$20 billion by 2026*<ul style="list-style-type: none">– Keppra market leader with approx. 2.2M patients & >€800M p.a.**• High proportion of refractory patients (¼ of new patients***) - combination treatments have limited therapeutic benefit• Large underserved patient population in need of improved treatment options
<h2>ADX71149: true synergistic MoA</h2>	<ul style="list-style-type: none">• Selective oral mGlu2 PAM with clear MoA in epilepsy• Showed 35-fold increase in Keppra efficacy in preclinical model• Potential first rational polypharmacy in epilepsy
<h2>Development path</h2>	<ul style="list-style-type: none">• Extensive preclinical and clinical data<ul style="list-style-type: none">– 8 Phase 1 and 2 Phase 2 studies• Janssen expect to start POC study in Q2 2021<ul style="list-style-type: none">– Top line data expected in H1 2022
<h2>Partnership with Janssen</h2>	<ul style="list-style-type: none">• Eligible to receive €109 million in pre-launch milestones and double digit royalties

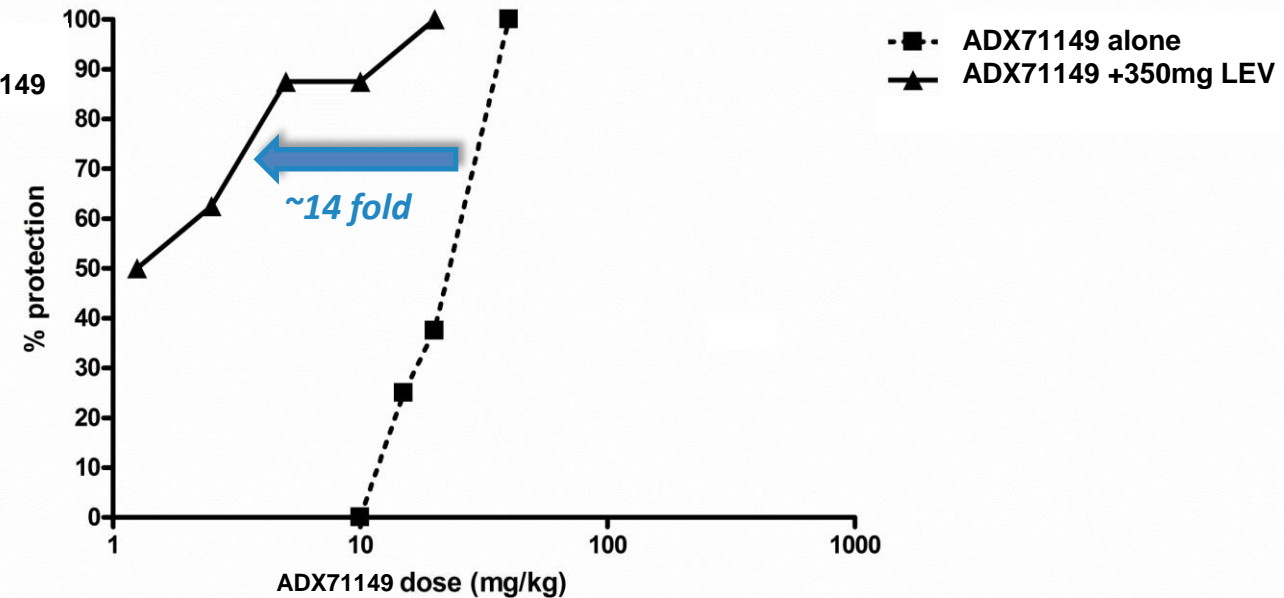
ADX71149 Preclinical Efficacy in Epilepsy - 6Hz Model

- Preclinical validation in pharmaco-resistant mouse epilepsy model:

ED₅₀ shift of Keppra by adding low dose of ADX71149

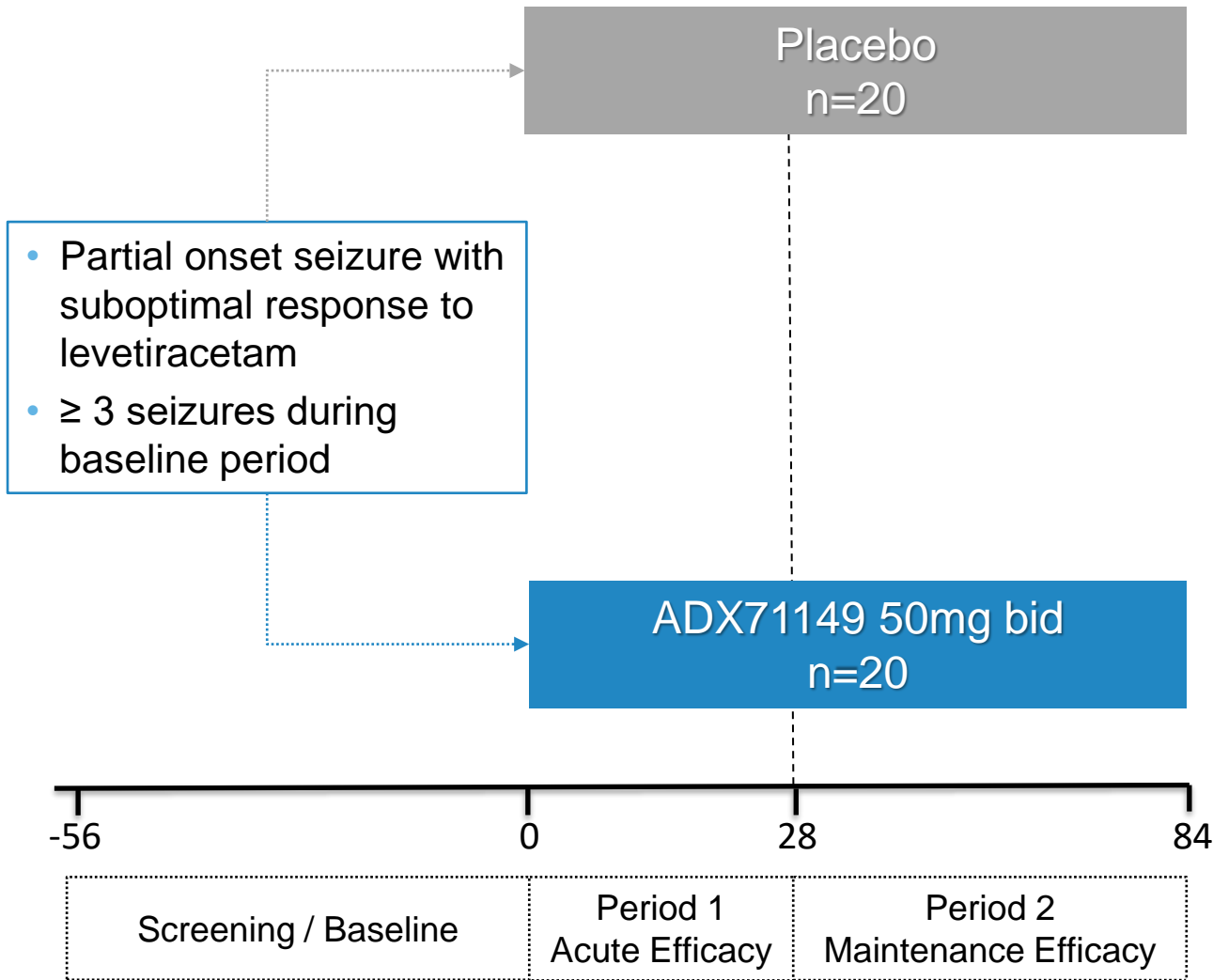


ED₅₀ shift of ADX71149 by adding ED₅₀ dose of LEV



- Keppra efficacy increased 35-fold when administered with a low dose of ADX71149
- Low dose of Keppra leads to 14-fold increase in efficacy of ADX71149
- True synergistic effect

ADX 71149 Phase 2a Epilepsy Study Design



- Double blind placebo controlled
- Establish 28-day seizure count (over 56-day baseline period)
- Primary endpoint: time to baseline seizure count
- Period 1: 4-week acute efficacy phase
- Period 2: 8-week maintenance efficacy phase
 - Subjects who do not reach or exceed their monthly baseline seizure count in Period 1 continue double-blind treatment during Period 2

Financials

Financials and Stock

- Cash runway through 2022
 - Cash at 31 Dec 2020: CHF18.7 million
 - Completed \$11.5M capital increase 11 Jan 2021
- No debt
- Traded on SIX Swiss Exchange: ADXN (ISIN:CH0029850754)
- ADS representing 6 shares traded on Nasdaq: ADXN (ISIN: US00654J107; CUSIP: 00654J107)
- 39.7M shares outstanding 56.2M (fully diluted, including treasury shares)
 - New Enterprise Associated - 14.21%
 - New Leaf Venture Partners - 4.77%
 - CAM Capital - 4.02%
 - Credit Suisse Asset Management - 3.01%
 - Management & board holds -12.15% (fully diluted basis)
- Analyst coverage:
 - Van Leeuwenhoek - Marcel Wijma
 - valuationLab - Bob Pooler
 - ZKB - Dr. Michael Nawrath
 - Baader Helvea AG - Bruno Bulic

Summary

3 clinical programs starting in H1 2021

- Dipraglurant PD-L1D registration study
- Dipraglurant blepharospasm Phase 2
- ADX71449 (J&J) epilepsy Phase 2

Technology and capabilities to deliver

- Experienced team of drug developers
- Pioneering allosteric modulation drug development
 - Proprietary screening assays and unique chemical library
- All programs developed in-house, protected with >200 patents

Solid foundation

- Partnerships with industry leaders
- Top tier US investors - NEA, NLV and CAM Capital Program
- Dual listed SIX Swiss exchange & US Nasdaq

Promising outlook

- Rich news flow in 2021 and beyond
 - Clinical programs
 - Multiple drug candidates in CCS



ALLOSTERIC MODULATORS FOR HUMAN HEALTH

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