

Innovative Treatments for Central Nervous System Disorders

February 2021

Allosteric modulators for human health

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Addex Overview

3 clinical programs	 Dipraglurant PD-LID 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	 Allosteric modulation is a validated & differentiated pharmacological approach to address drug targets Proprietary biological screening assays and chemical library
In house discovered pipeline	 Multiple novel drug candidates entering clinical candidate selection Driving long term growth & future partnership opportunities
Partnerships	 J&J - €109M in milestones & double digit royalties Indivior - \$330M in milestones, royalties up to double digit & funded research program
Top tier US investors	 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



* pending removal of governmental and institutional restrictions and lessening of the impact of the global coronavirus pandemic on the U.S. healthcare system, which delayed our previously anticipated initiation in the first quarter of 2020

Addex Pipeline

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

Lead Program Entering US Pivotal Study



NAM = Negative Allosteric Modulator PAM = Positive Allosteric Modulator

Experienced Team

Leadership Team			
Tim Dyer CEO / CFO	Dr Roger Mills Chief Medical Officer	Dr Robert Lutjens Head of Discovery Biology	Dr Jean-Philippe Rocher Head of Discovery Chemistry
Co-Founder of Addex Formerly with PwC UK Chartered Accountant	Developed Nuplazid in PD Psychosis 30 years in Pharma industry including Pfizer, Gilead and Acadia	Member of Addex founding team Formerly with Glaxo & Scripps Research Institute	Member of Addex founding team Formerly with Pierre Fabre, GSK and Mitsubishi
Non-executive Directors			
Vincent Lawton Chairman	Ray Hill Board member	Jake Nunn Board member	Isaac Manke Board member
Former European Head of Merck & Co. Former MHRA Board member	Former Executive Director Merck & Co.	Venture advisor and former Partner at New Enterprise Associates	General Partner at Acorn Bioventures. Formerly Partner at New Leaf Venture Partners
Scientific Advisory Board			
Darryle Schoepp Chairman of SAB	Mark Bear Picower Prof. of Neuroscience at MIT	Peter Bernstein Principal, PhaRmaB LLC	Benny Bettler Biomedicine Prof. at Basel University
Former leader of Neuroscience research department at Eli Lilly, and at Merck was Neuroscience research therapeutic area leader	Formerly on faculty of Brown University School of Medicine and an Investigator of the Howard Hughes Medical Institute	Formerly with ICI Astra Zeneca Awarded numerous accolades including Fellow of the American Chemical Society	Formerly at Novartis and discovered allosteric modulators at GABA _B receptor and recipient of the Peter Speiser Award



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

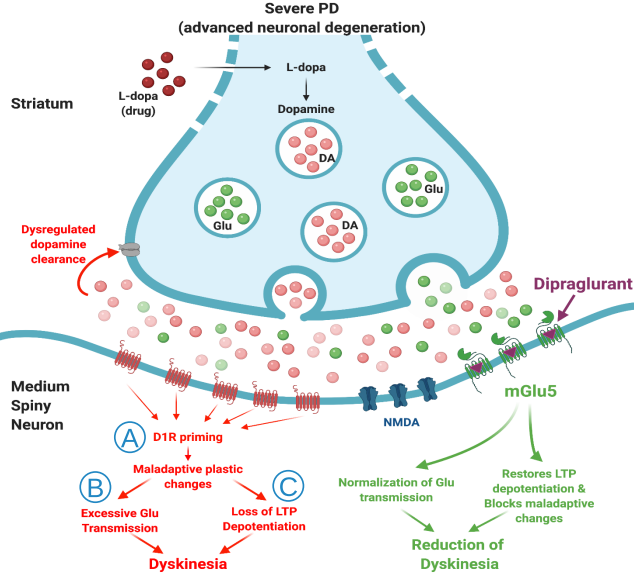


Disability and Impact of PD-LID

Invariably associated with	 Dyskinesias caused by neurodegeneration Depending replacement lowers the triggering threshold for symptoms
long-term L-dopa use	 Dopamine replacement lowers the triggering threshold for symptoms LID can become as disabling as the PD symptoms themselves
	Uncontrollable muscle contractions, twisting and writhing
Symptoms include dystonia,	 Painful and severely disabling
chorea, and choreoathetosis	 Causes fatigue/exhaustion and increased risk for falls and injuries
	 Social withdrawal, reduced quality of life and increased burden on caregiver
Drovolopoo rolatad ta diagooo	 >40% of patients experience LID within 4-6 years of L-dopa treatment
Prevalence related to disease duration	 Increases to 90% after 9 -15 years
Guration	 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

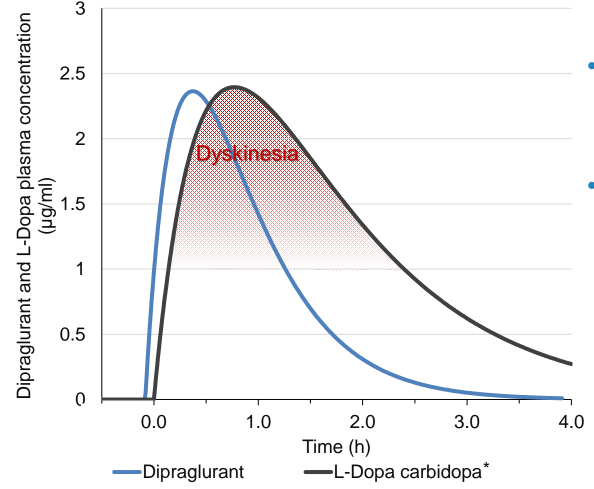


- Loss of substantia nigra neurons combined with the non-physiological, pulsatile stimulation of dopamine receptors with L-dopa are at the basis of LID development
- In the striatum, LID is the result of:
 - \bigcirc D1 receptor priming
 - B Excess glutamate transmission
 - C Loss of LTP depotentiation
- 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



LTP = Long Term Potentiation D1R = D1 dopamine receptor Glu = glutamate DA = dopamine

Dipraglurant PK is a Key Advantage for Treating LID



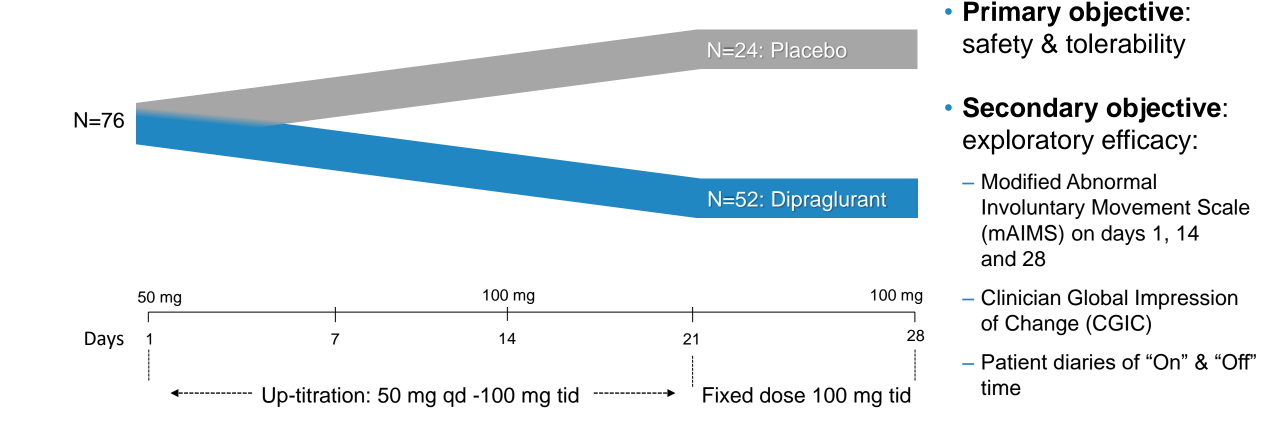
- Dyskinesia symptoms are correlated to peak levels of L-dopa
- PK of dipraglurant allows control of glutamatergic tone ahead of L-dopa Cmax

Dipraglurant normalizes abnormal glutamate stimulation during peak levodopa dose

Dipraglurant peaks ahead of L-dopa for optimal LID control



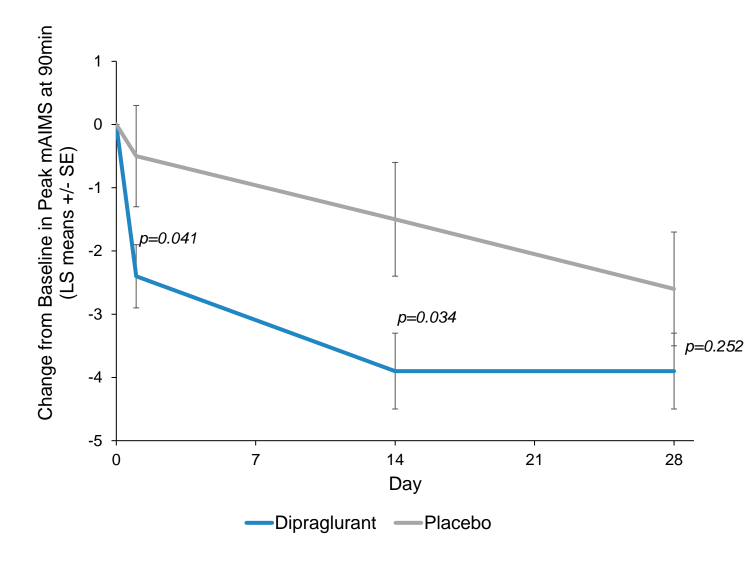
Dipraglurant Phase 2a Study in LID (in US and Europe)



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



Dipraglurant Improves LID by 30%

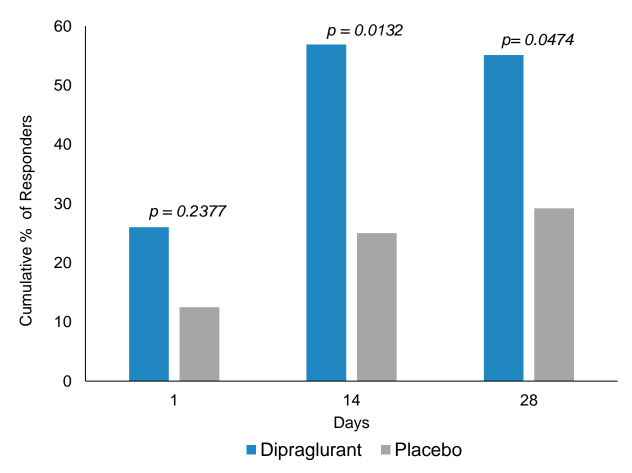


- Statistically significant effects: Day 1 (50mg) and Day 14 (100mg)
- Improvement maintained through Day 28
- Increasing placebo response caused significance to be lost at Day 28
- No placebo mitigation in study

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 ≥ 30% improvement on mAIMS



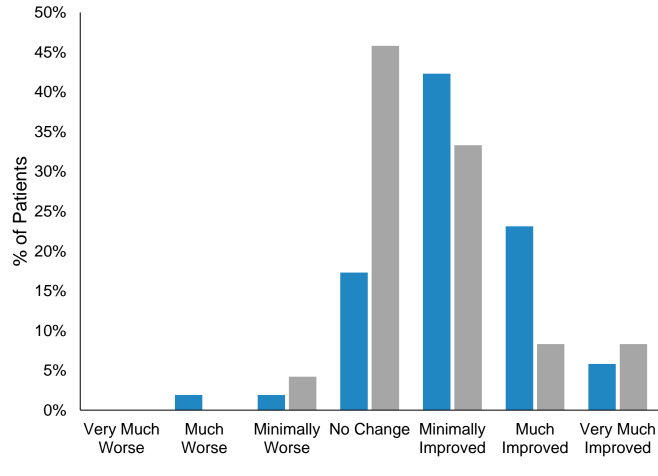
Responder analysis (≥30% change of mAIMS from baseline)				
Midday dose	Dipraglurant Placebo		acebo	
Day 1 (50 mg)	n=13	26.0%	n=3	12.5%
Day 14 (100 mg)	n=29	56.9%*	n=6	25.0%
Day 28 (100 mg)	n=27	55.1%*	n=7	29.2%

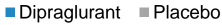
*statistically significant

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
	 Recommended by Movement Disorder Society (MDS) 	 Suboptimal for detecting treatment-related changes
	 FDA regulatory precedent (GOCOVRI® approval) 	 Limited to patient assessments
Characteristics	 Contains anchored objective clinician evaluated measures of dyskinesia 	 Prone to placebo effect
	 Includes both patient and physician assessments of impairment 	
	 Less prone to placebo effect 	
Clinimetrics	 Validated 	 Only the original version has been validated
Development	 Developed in 2009 specifically for dyskinesia in PD 	 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

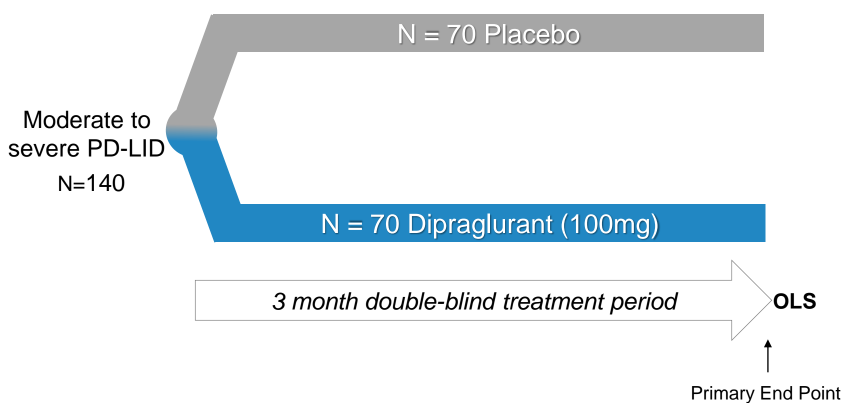


Dipraglurant PD-LID Registration Program Started

- 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



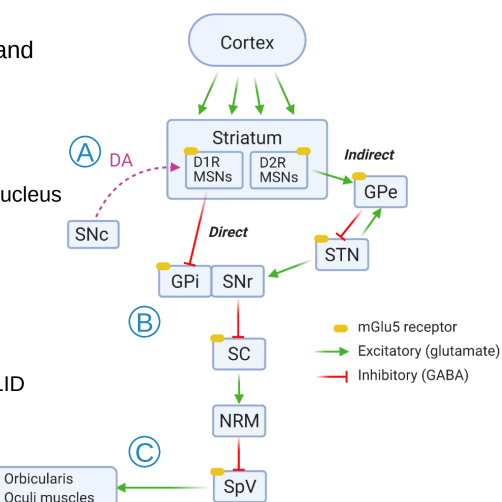
Blepharospasm (BSP)

- 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

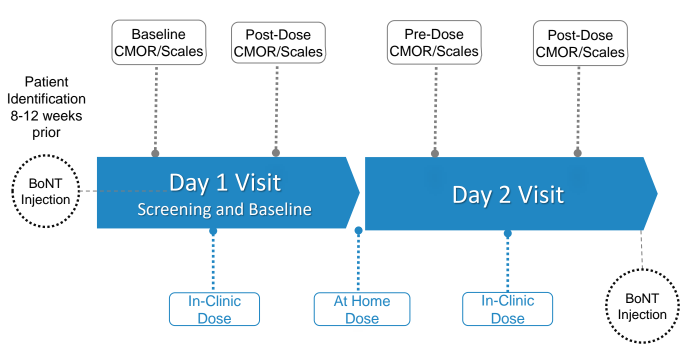
- Dystonias are neuro-functional rather than neuro-degenerative
- Common features include alterations in neuronal connectivity/function and synaptic communication
- BSP pathophysiology is linked to:
 - A Reduction of dopamine input into striatum
 - B Increased inhibition of direct pathway from superior colliculus to trigeminal nucleus
 - C 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)



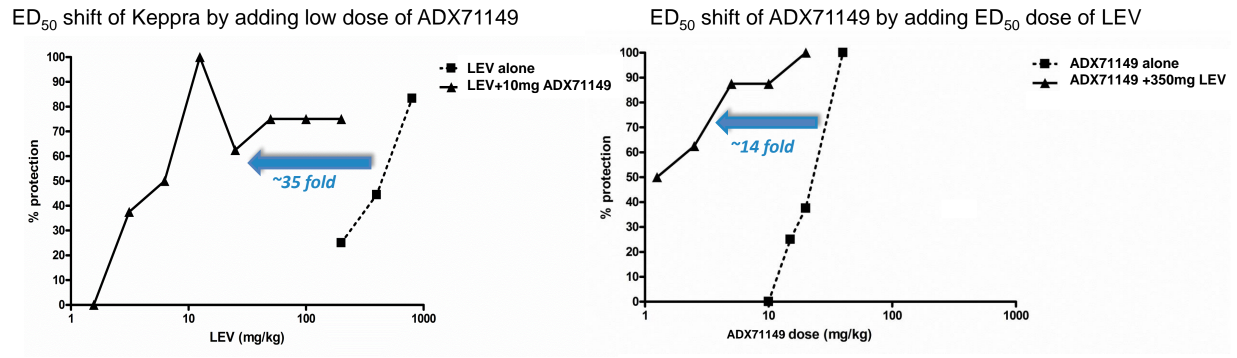
ADX71149 Opportunity in Epilepsy

	 Market projected to reach \$20 billion by 2026*
Large market & unmet medical need	– 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
	 Selective oral mGlu2 PAM with clear MoA in epilepsy
ADX71149: true	 Showed 35-fold increase in Keppra efficacy in preclinical model
synergistic MoA	 Potential first rational polypharmacy in epilepsy
	 Extensive preclinical and clinical data
Dovelopment noth	 8 Phase 1 and 2 Phase 2 studies
Development path	 Janssen expect to start POC study in Q2 2021
	 Top line data expected in H1 2022
Partnership with Janssen	 Eligible to receive €109 million in pre-launch milestones and double digit royalties



ADX71149 Preclinical Efficacy in Epilepsy - 6Hz Model

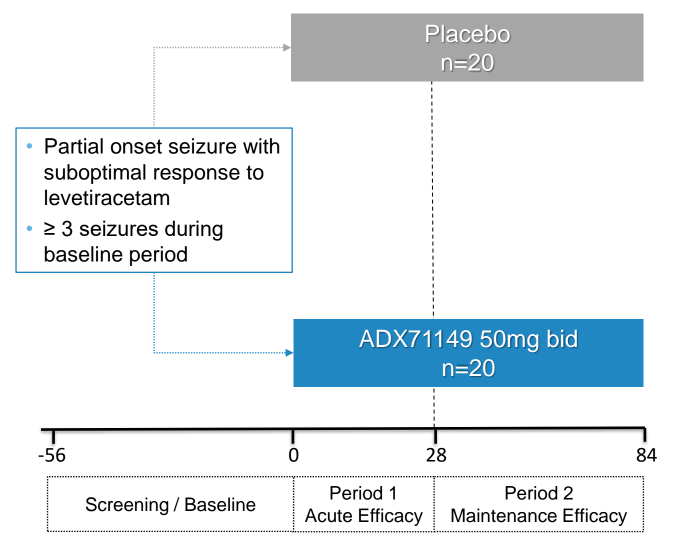
• Preclinical validation in pharmaco-resistant mouse epilepsy model:



- 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

2 divided programs starting	 Dipraglurant PD-LID registration study
3 clinical programs starting in H1 2021	 Dipraglurant blepharospasm Phase 2
	 ADX71449 (J&J) epilepsy Phase 2
	 Experienced team of drug developers
Technology and capabilities	 Pioneering allosteric modulation drug development
to deliver	 Proprietary screening assays and unique chemical library
	 All programs developed in-house, protected with >200 patents
	 Partnerships with industry leaders
Solid foundation	 Top tier US investors - NEA, NLV and CAM Capital Program
	 Dual listed SIX Swiss exchange & US Nasdaq
	 Rich news flow in 2021 and beyond
Promising outlook	 Clinical programs
	 Multiple drug candidates in CCS





ALLOSTERIC MODULATORS FOR HUMAN HEALTH

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