



IMPROVING THE LIVES OF PEOPLE WITH NEURODEVELOPMENTAL DISABILITIES

17 November 2022

Jefferies London Healthcare Conference 2022

Forward looking statements

This presentation contains forward looking statements that involve risks and uncertainties. Although we believe that the expectations reflected in the forward looking statements are reasonable at this time, Neuren can give no assurance that these expectations will prove to be correct. Actual results could differ materially from those anticipated. Reasons may include risks associated with drug development and manufacture, risks inherent in the regulatory processes, delays in clinical trials, risks associated with patent protection, future capital needs or other general risks or factors.

1

NDA for trofinetide to treat Rett syndrome accepted by FDA for Priority Review with PDUFA date set for 12 March 2023

2

Potential revenue from Acadia in 2023 for Rett syndrome in the US alone of US\$73m (A\$112 million)¹ plus double-digit % royalties

3

Strong partnering interest received for trofinetide outside North America

4

Accelerating Phase 2 development of NNZ-2591 in 4 indications, with potential markets 5x Rett syndrome

5

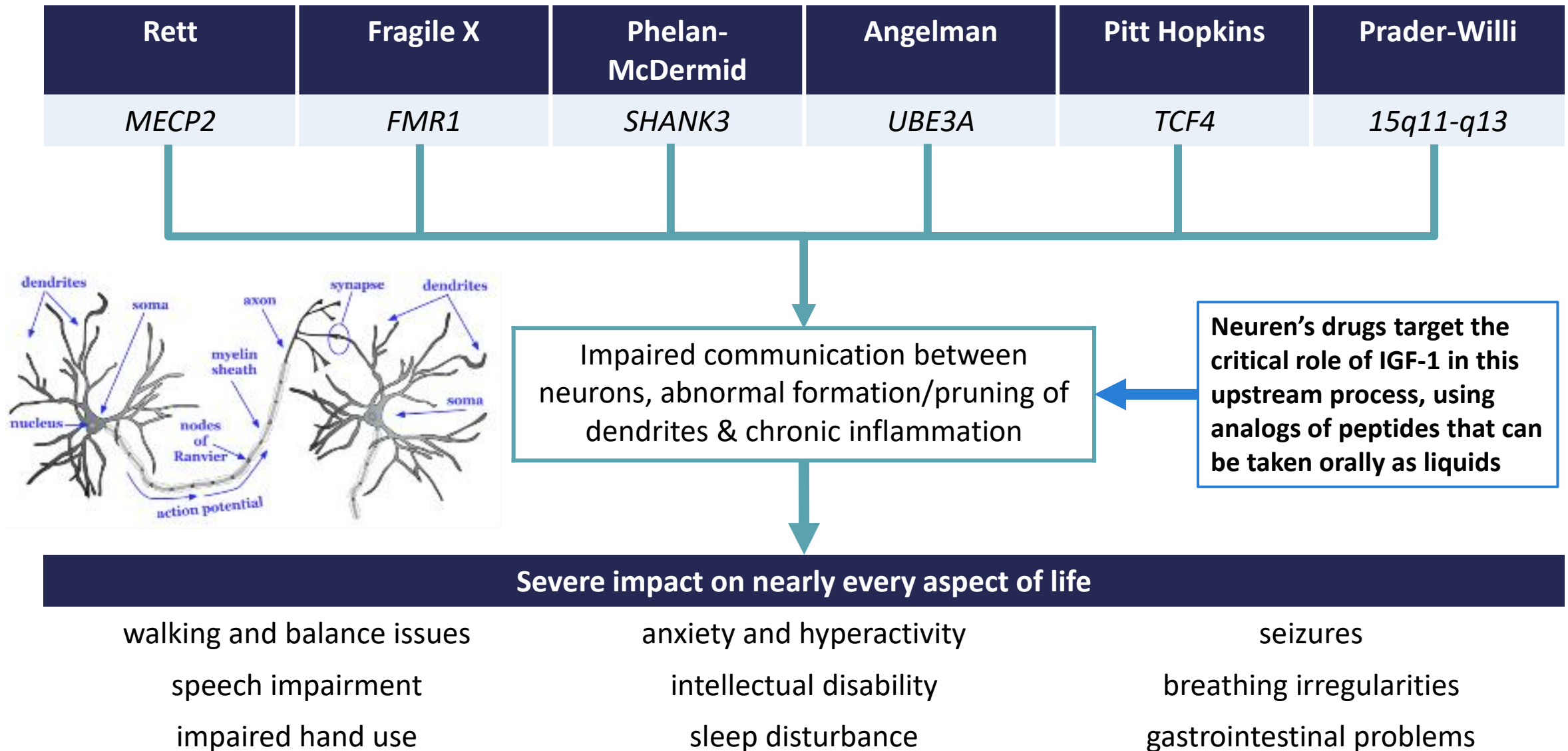
NNZ-2591 novel mechanism of action has many more potential applications

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


A\$27 million cash at 30 Sep 2022, plus US\$10m receipt in Oct – well funded to execute NNZ-2591 Phase 2 trials and preparation for Phase 3

¹ Assuming a New Drug Application (NDA) is approved by the FDA, the product is launched in the US, US\$33m is received as one third share of the value of a Rare Pediatric Disease Priority Review Voucher if awarded upon approval of a NDA, and a USD/AUD exchange rate of 0.65

Seeking a ground-breaking impact on neurodevelopmental disorders



All development programs at Phase 2 or later

Compound	Indication	Preclinical	Phase 1	Phase 2	Phase 3	Registration	Commercial rights
Trofinetide	Rett ²					US PDUFA date 12 Mar 2023 ¹	NA:  ACADIA
	Fragile X ²						RoW: 
NNZ-2591	Phelan-McDermid ³				Results H1 2023		
	Angelman ³						
	Pitt Hopkins ³						
	Prader-Willi ⁴	Results H2 2023					

¹ Priority Review granted

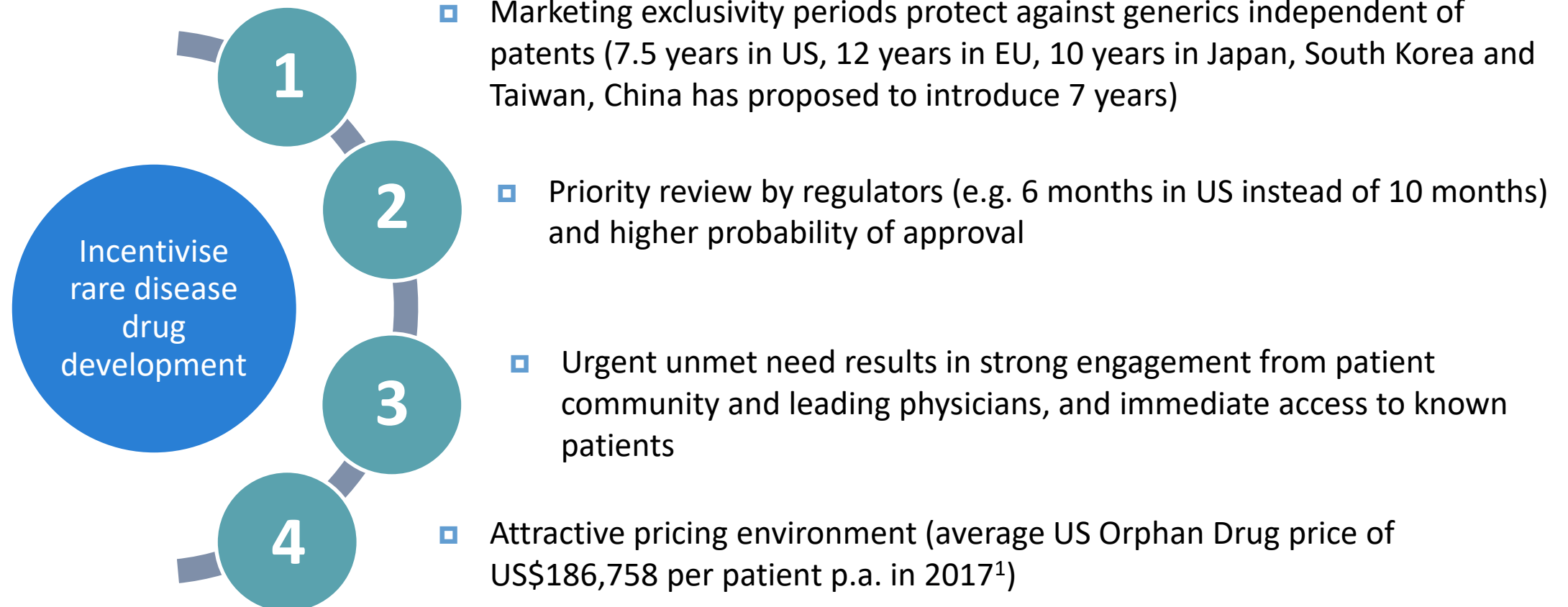
² Orphan Drug designation in US and EU, Fast Track designation in US

³ Orphan Drug designation in US and EU

⁴ Orphan Drug designation in US

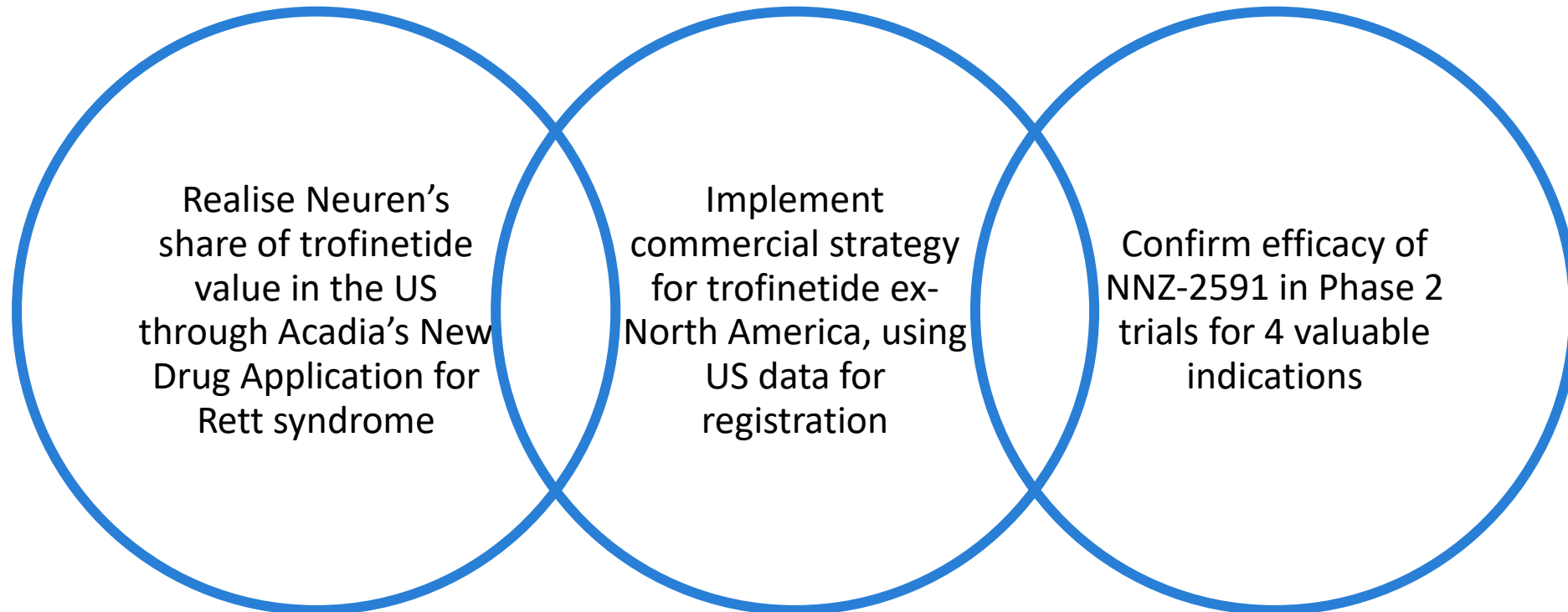
Attractiveness of Orphan Drug model

Neuren is targeting multiple “rare diseases”, but they are not “ultra-rare”



¹ AHIP

Three key drivers transforming near term value

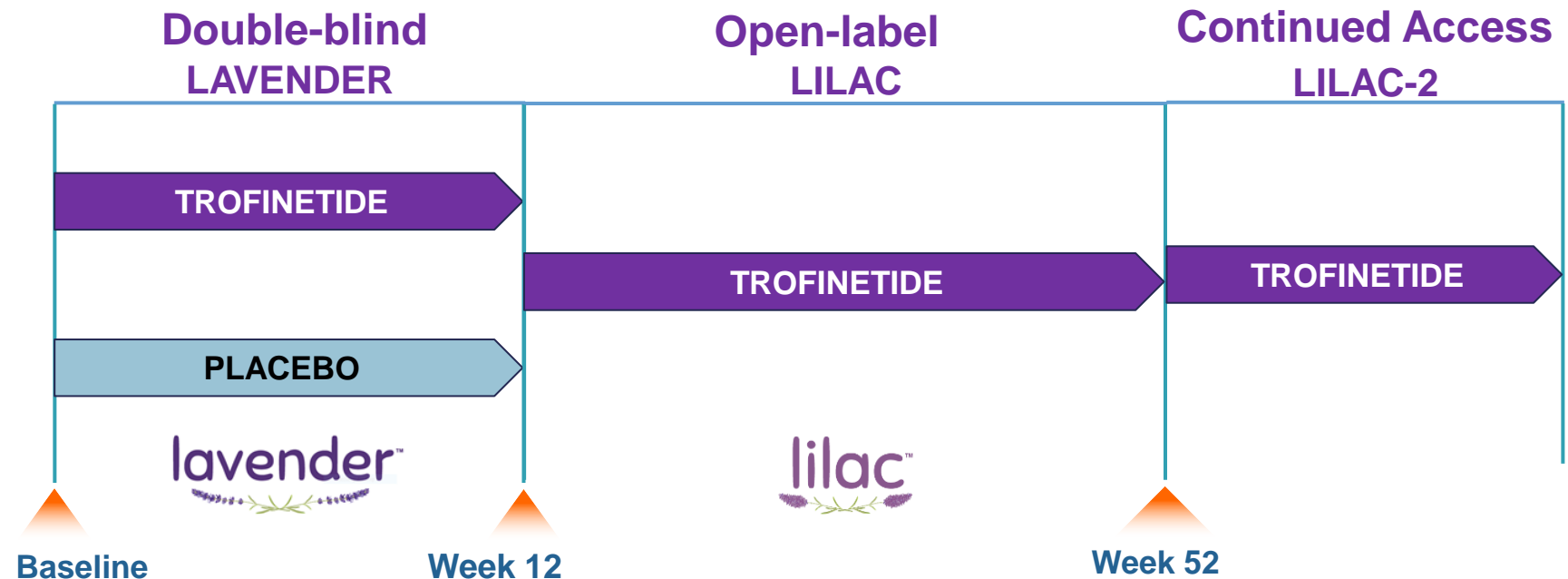


Rett syndrome Phase 3 and NDA

- Acadia submitted NDA in July 2022 for treatment of Rett syndrome in patients two years of age and older
- NDA based on pivotal efficacy from positive Phase 3 trial, supportive efficacy from Neuren's positive Phase 2 trial, safety data from completed and ongoing studies
- FDA accepted NDA for Priority Review - PDUFA action date set for 12 March 2023
- FDA advised that at this time it is not planning to hold an Advisory Committee meeting

LAVENDER™ randomised, double-blind, placebo-controlled trial:

- 187 females aged 5 to 20 years
- RSBQ (caregiver) and CGI-I (physician) at 12 weeks co-primary efficacy endpoints



North America

- ## Ex-North America

- 9

5x larger opportunity for NNZ-2591

Disorder	Gene mutation	Published prevalence estimates	Potential patients		
			US ¹	Europe ¹	Asia ^{1, 2}
Phelan-McDermid	<i>SHANK3</i>	1/8,000 to 1/15,000 males and females	22,000	28,000	81,000
Angelman	<i>UBE3A</i>	1/12,000 to 1/24,000 males and females	14,000	18,000	52,000
Pitt Hopkins	<i>TCF4</i>	1/34,000 to 1/41,000 males and females	7,000	9,000	25,000
Prader-Willi	<i>15q11-q13</i>	1/10,000 to 1/30,000 males and females	13,000	16,000	47,000
			56,000	71,000	205,000

- Current opportunity for NNZ-2591 is more than 5 times the Rett Syndrome opportunity³
- There are many other neurodevelopmental disorders potentially relevant for NNZ-2591 mechanism of action
- Neuren retains global rights

¹ Estimates derived by applying the mid-point of the prevalence estimate range to the populations under 60 years

² Asia comprises Japan, Korea, Taiwan, Israel and urban populations of China and Russia

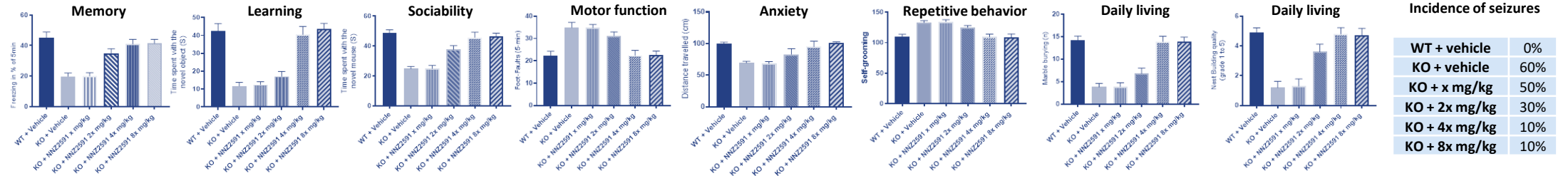
³ Based on number of potential patients globally

NNZ-2591 has ideal attributes leading into Phase 2

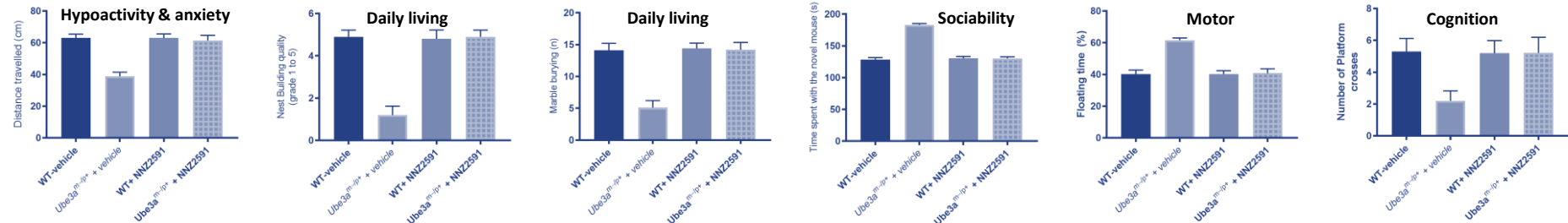
- ✓ Novel mechanism of action
- ✓ Clear and consistent efficacy in mouse models of each syndrome
- ✓ Biochemical effects in the brain confirmed
- ✓ Optimum dose identified
- ✓ Demonstrated high oral bioavailability and blood-brain barrier penetration
- ✓ IND-enabling program of non-clinical toxicology and CMC studies completed
- ✓ Proprietary drug substance manufacturing process with exceptional purity and high yield, administered as patient-friendly liquid dose
- ✓ Safe and well tolerated in Phase 1 trial
- ✓ Orphan designations from FDA and EMA

Clear and consistent efficacy in animal models

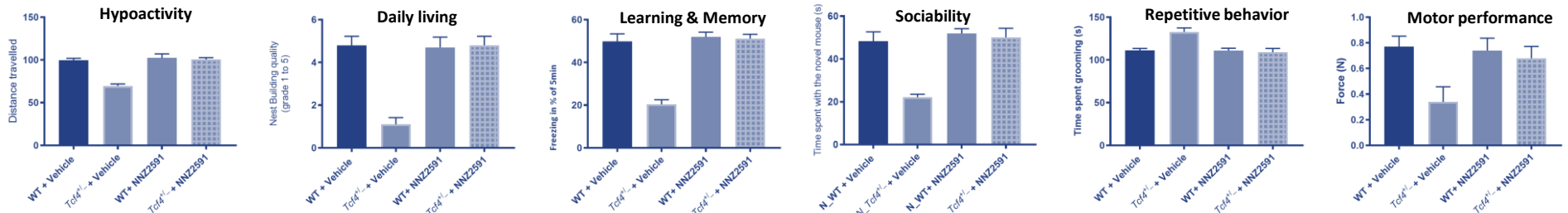
Phelan-McDermid



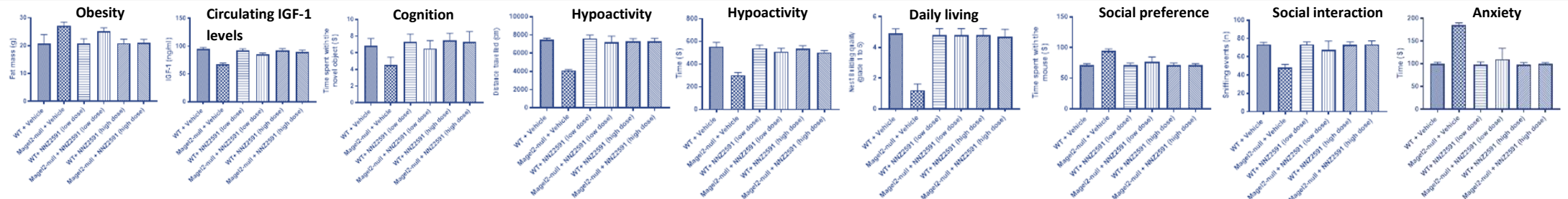
Angelman



Pitt Hopkins

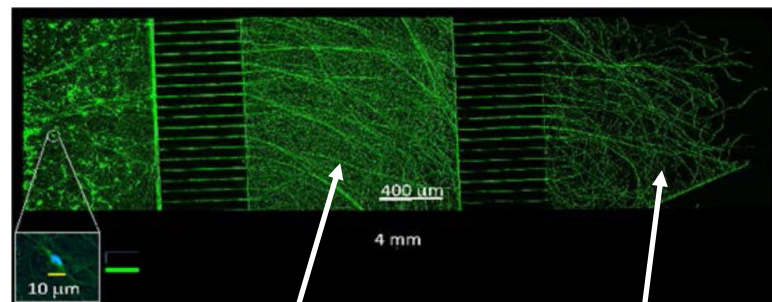
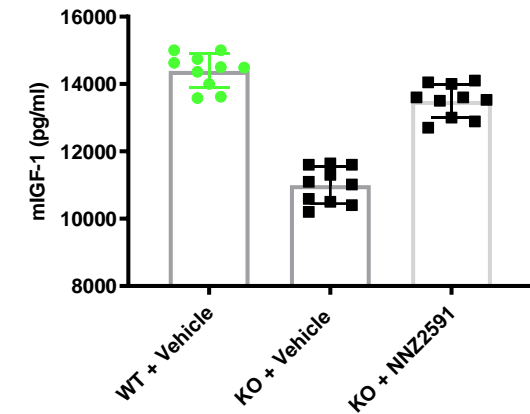
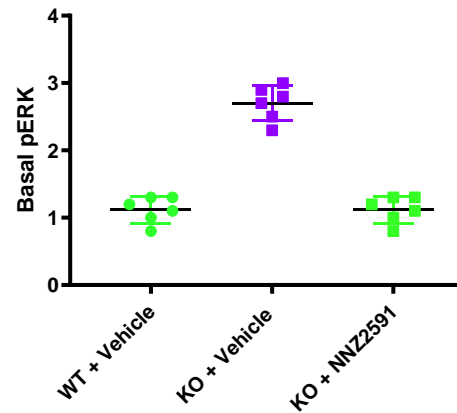
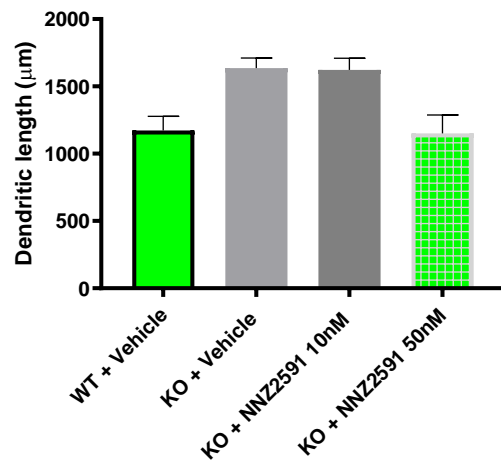


Prader-Willi



Biochemical effects confirmed

In biochemical testing, NNZ-2591 was shown to normalise the abnormal length of dendrite spines between brain cells, the excess activated ERK protein (pERK) and the depressed level of IGF-1 in *shank3* knockout mice



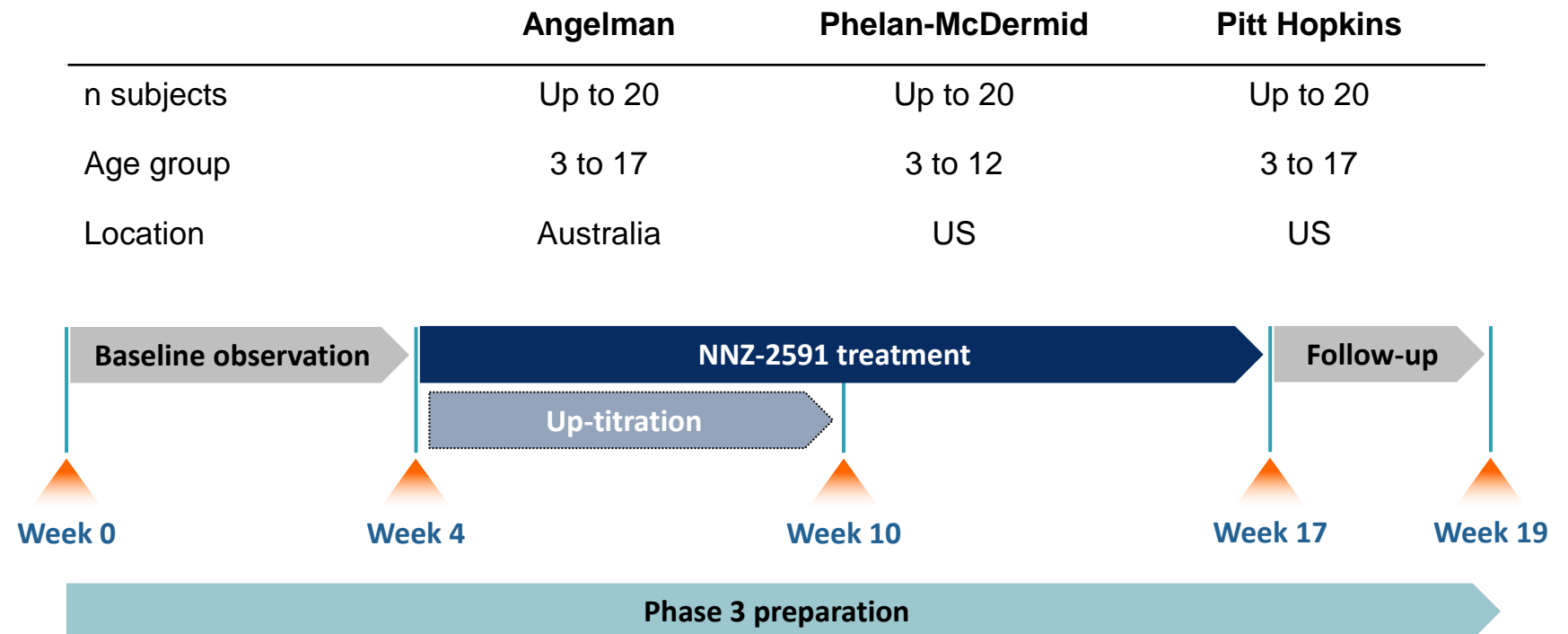
Abnormal dendrites in
shank3 knockout mice

Normalisation after treatment
with NNZ-2591

Key features of first Phase 2 trials

Overall aim – expedite data that enables subsequent trials to be designed as registration trials and prepare for Phase 3 in parallel

- Prioritising speed to data
- Maximising opportunity to demonstrate effects
- Confirm safety and PK in pediatric patients
- Assess treatment impact across multiple efficacy measures to select primary endpoint for registration trial



On track to deliver significant value upside over next 18 months

Prader-Willi syndrome Phase 2 trial results (H2 2023)

Phase 2 trial results in Angelman, Phelan-McDermid and Pitt Hopkins syndromes (H1 2023)

Approval of NDA for Rett syndrome (Q1 2023)

Commercial partnerships ex-North America for Rett syndrome

Commence Prader-Willi syndrome Phase 2 trial (file IND Q4 2022)

✓ **FDA acceptance of NDA filing for Rett syndrome**

✓ **Commence Phelan-McDermid and Pitt Hopkins syndromes Phase 2 trials**

✓ **Acadia submits New Drug Application (NDA) for Rett syndrome**

✓ **Commence Phase 2 trial in Angelman syndrome**

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