Dose Responsiveness of LUM-201 as Measured by Acute GH Response and IGF-1 and Annualized Height Velocity (AHV) Measured at 6 Months in the Interim Analysis of the OraGrowtH212 Study in Idiopathic Pediatric Growth Hormone Deficiency (iPGHD).

Cassorla F<sup>1</sup>, MD; Román R<sup>1</sup>, MD; Johnson M<sup>2</sup>, PhD; Smith C<sup>2</sup>, MS; Avila A<sup>1</sup>, RN; Iñiguez G<sup>1</sup>, PhD; Baier I<sup>1</sup>, MD; Said D<sup>1</sup>, RN; Karpf DB<sup>2</sup>, MD; McKew JC<sup>2</sup>, PhD; Thorner M<sup>2</sup>, MB BS, DSc



<sup>1</sup>University of Chile, Santiago, Chile <sup>2</sup>Lumos Pharma, Austin, TX

#### Disclosure

Dr. Cassorla is an investigator for clinical studies with LUM-201 at the University of Chile (Sponsor - Lumos Pharma, Inc.) and has previously acted as a consultant for Debiopharm, Pfizer, Merck, Novo Nordisk and Sandoz.

LUM-201 is an investigational compound and is not approved for use by the FDA or any other regulatory agency. Some of the slides in this presentation are derived or copied from corporate presentations previously given by Lumos Pharma, Inc. These slides are used with permission.



## LUM-201 (ibutamoren) – Mechanism of Action



## Oral LUM-201 is a *growth hormone (GH)* secretagogue

- Acts as a durable agonist of GH Secretagogue Receptor (GHSR1a) to stimulate GH release<sup>1</sup>
- LUM-201 has been observed to increase the amplitude of endogenous, pulsatile GH secretion over 24 hours<sup>2,3</sup>
- Another differentiating feature vs rhGH is the *natural negative feedback mechanisms, which limit the potential for hyperstimulation and excessive increases in IGF-1*
- LUM-201 promotes pulsatile GH secretion in a selective PGHD Population



Moderate Idiopathic PGHD - Axis Responsive



- 1. Howard 1996 Science 273:974-977
- 2. Nass 2008 Ann Intern Med 149:601-611
- 3. Chapman 1997 J Clin Endocrinol Metab 82:3455-3463

#### Phase 2- Pulsatility and PK/PD Study Design Naive Idiopathic PGHD Patients



#### **Study Information**

- Open-label study: N = 22
- Pre-pubertal PGHD subjects that are rhGH-treatment naïve
- Inclusion: Height < 2 SD, delayed bone age, peak GH response to a clonidine stimulation test between 3 and 10 ng
- Dosing to near-adult height
- Single, specialized clinical site University of Chile, Santiago

#### **Primary Endpoints:**

 Assess LUM-201 effect on endogenous GH pulsatility and Annualized Height Velocity (AHV)

OraGrowtH212

• Evaluate PK/PD in children

#### Goals:

- Confirm prior PK/PD data in adults & subset of Merck 020 trial
- Support future regulatory filings & commercialization



1. Does LUM-201 dose-dependently augment endogenous GH pulses in patients with Idiopathic Pediatric Growth Hormone Deficiency (iPGHD)?

2. Will increased amplitude of GH pulsatility and increase in IGF-1 within normal range improve height velocity?

3. Is the effect on AHV durable out to 12 months?



#### **Baseline Demographics**

6

Subjects N=15	1.6 mg N=8	3.2 mg N=7
	Mean (SD)	
Age (mos)	96.9 (11.9)	<b>95.0</b> (22.7)
Height (cm)	115.2 (4.57)	113.1(9.97)
Height SDS	-2.12 (0.29)	-2.34 (0.45)
IGF-1 SDS	-1.1 (0.535)	-0.8 (0.377)
MPH (cm)	161.8 (6.98)	160.82 (5.73)
MPH SDS $\Delta$	0.73 (0.47)	0.81 (0.43)
BA Delay (yrs)	1.50 (0.26)	1.83 (0.88)
BMI (SDS)	-0.18 (0.96)	+0.48 (1.02)
Male/Female%	63/37	<b>71</b> /29



# Differences between the two groups:

- Slight imbalance in age and gender
- Slight imbalance in delta below MPH, BMI, and bone age delay



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## AHV Before and After 6 months of LUM-201 Treatment



# 

#### 6-month observations:

- LUM-201 raised the AHV (growth rate) from baseline after 6 months on therapy for both the 1.6 mg/kg cohort (p = 0.0006) and the 3.2 mg/kg cohort (p < 0.0001)</li>
- No statistical difference exists between the two cohorts at each timepoint
- As expected, greater growth response was observed in patients with lower baseline height velocity



# Durable Response After 12 Months of LUM-201 Administration







IGF-1 Values: Treatment with LUM-201 Increased Serum IGF-1 Concentration and IGF-1 SDS Values

IGF-1 IGF-1 SDS P = 0.0329P = 0.0228400-0.70 P = 0.0036P = 0.0048SDS 226.4 1-300-0.01 mean IGF-1 (ng/mL) 177.1 mean IGF-1 5 0 200 -8 112.3 119.0 -1-100 --0.80 -0.92 7 5 7 8 -2 0 3.2 1.6 3.2 1.6 1.6 3.2 1.6 3.2 baseline 6 mo baseline 6 mo



#### **Conclusions :**

- There is a significant increase in IGF-1 levels that remains within the normal range
- Based on the MOA of LUM-201, these data support the physiological IGF-1 feedback



## IGF-1, GH Pulsatility, Height Velocity: Patient A **1.6 mg**/kg/day





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\*\*Percent change from baseline calculated as: (6mo value – baseline value) / (baseline value)

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#### Interim Analysis Safety Profile

#### **Safety Profile:**

- No treatment-related Serious Adverse Events (SAEs) or Severe AEs
- No meaningful safety signals observed in either laboratory values, adverse event data, or in electrocardiogram values.

#### Most Common AEs (% of subjects) noted are:

- Transient increased appetite (76.5%)
- Pain in extremity (17.6%)
- Arthralgia (11.8%)
- Abdominal pain (5.9%)
- Influenza (5.9%)

#### **Safety Conclusion:**

 At time of interim analysis, LUM-201 was well tolerated and showed no significant safety signals







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- Based on Interim Analysis data, OraGrowtH212 data demonstrates that growth acceleration is durable through 12 months in our study population, pre-pubertal, treatment naïve idiopathic PGHD patients.
- No statistical difference exists between the cohorts at any time point.
- Due to some baseline imbalance, the optimal dose cannot be determined from this data set.
- We plan to continue the OraGrowtH212 Trial until near adult height.
- The observed growth is in line with rhGH historical growth of 8.3-8.6 cm (KIGS <sup>1</sup>, GeNeSiS <sup>2</sup>) in this moderate idiopathic pre-pubertal PGHD population.



Sources: <sup>1</sup> Blum et al JES 2021, <sup>2</sup> Ranke et al JCEM 2010

#### University of Chile, Santiago Institute of Maternal and Child Research Pediatric Team

## OraGrowtH212



