Continued Efficacy and Safety with up to 2 Years of Treatment with Lonapegsomatropin (TransCon[™] hGH) in Children with Growth Hormone Deficiency

Donna Campbell¹, Kimberly Walsh², Natalie Marlen³, Wenjie Song⁴, Meng Mao⁴, Steven Chessler⁴, Allison Komirenko⁴, Michael Beckert⁴, Aimee D. Shu⁴

'Rocky Mountain Pediatric Endocrinology, Centennial, CO, USA; "Children's Hospital at Dartmouth-Hitchcock, Lebanon, NH; "Center of Excellence in Diabetes and Endocrinology, Sacramento, CA; ⁴Ascendis Pharma, Inc, Palo Alto, CA

BACKGROUND

 Once-weekly lonapegsomatropin (TransCon hGH) is an investigational prodrug for growth hormone deficiency (GHD) that consists of 3 components: unmodified somatropin, an inert carrier that protects it, and a linker that temporarily ioins the two1.2 (Figure 1)

Figure 1, Lonapegsomatropin (TransCon hGH) Design



Tissue Distribution Physiological levels Therapeutic effects: efficacy, safety and tolerability

· In the pivotal phase 3 heiGHtTrial evaluating treatment-naïve children with GHD, lonapegsomatropin demonstrated superior annualized height velocity (AHV) and statistically greater change from baseline in height standard deviation score (A height SDS) at 52 weeks compared to daily somatropin therapy (Genotropin) and had a similar safety and tolerability profile

· In the phase 3 fliGHtTrial, children who switched from daily somatropin to lonapegsomatropin continued to grow well and maintained a good safety profile4

· Results are reported from heiGHt and fliGHt subjects who continued into the phase 3 enliGHten open-label long-term extension (OLE) Trial for up to 52 weeks (data cut: June 1st 2020)

METHODS

TRIAL DESIGN

Phase 3 heiGHt Tria

 heiGHt was a 52-week, open-label, active-controlled, pivotal phase 3 trial in which treatment-naïve, prepubertal subjects (males 3-12; females 3-11 years old) with GHD were randomized 2:1 to receive once-weekly lonapegsomatropin 0.24 mg hGH/kg/week via vial/svringe or an equivalent weekly dose of daily Genotropin via pen device

Phase 3 fliGHt Trial

· fliGHt was a 26-week, open-label phase 3 trial in which treatment-experienced subjects (6 months to 17 years old; subjects < 3 years old could be treatment-naïve) with GHD switched from their previous daily somatropin to lonapegsomatropin 0.24 mg hGH/kg/week via vial/syringe

Phase 3 enliGHten OLE Trial

· All subjects who enrolled into the long-term extension trial received lonapegsomatropin at their previous dose via vial/syringe (Genotropin subjects from heiGHt started lonapegsomatropin 0.24 mg hGH/kg/week) (Figure 2) · Subjects in the US switched to the TransCon hGH Auto-Injector when available

Figure 2. Lonapegsomatropin Phase 3 Clinical Program



OUTCOMES

- Efficacy · Growth outcomes were evaluated approximately every 13 weeks
- · Three groups were analyzed:
- Treatment-naive subjects treated with
- Ionapegsomatropin in heiGHt, followed by continuation of lonapegsomatropin in enliGHten
- Treatment-naive subjects treated with daily somatropin in heiGHt, followed by lonapegsomatropin in enliGHten
- Subjects previously treated with daily somatropin who switched to lonapegsomatropin in fliGHt, followed by continuation of lonapegsomatropin in enliGHten

Disposition, Demographics, and Baseline Characteristics

subjects from fliGHt at Week 78

The mean dose of lonapegsomatropin remained

Table 2. Demographics and Baseline Characteristics

8.5 (2.7)

3.3, 13.1

86 (81.9)

-2.3 (1.1)

-2.1 (0.9)

5.9 (2.8)

37 (35.2)

approximately 0.24 mg hGH/kg/wk for subjects from

heiGHt at Week 104 and was 0.20 mg hGH/kg/wk for

Age

Mean (SD)

Min, max

Gender, Males, n (%)

Height, Mean (SD)

Mean (SD)

≤5 ng/mL, n (%)

>5 ng/mL, n (%)

Height SDS, Mean (SD)

IGF-1 SDS, Mean (SD)

Peak stimulated GH level prior to initia

Nearly all subjects who completed heiGHt (158/159) Table 1. Subject Dispositio

	and fliGHt (140/144) continued into enliGHten					
	(Table 1)		heiGl	łt	fliGHt	
	 Eight (2.7%) subjects have prematurely withdrawn from the trial for the following 		Lonapegaomatropin n (%)	Daily somstropin n (%)	Lonapagaomatroj n (%)	
	reasons: 4 subjects (1.3%) due to withdrawn consent, 2 (0.7%) for protocol violation, and 2 for "other" reasons	Enrolled and dosed in parent trial	105	56	146	
	 As of the data cut, 2 subjects have achieved near adult height (AHV <2 cm/year over the last 9 months or bone age >14 [females] or >16 	Completed parent trial	104 (99.0)	55 (98.2)	144 (98.6)	
•	[males]) and thus have completed the trial Baseline demographics were balanced between	Enrolled and dosed in enliGHten	103 (98.1)	55 (98.2)	140 (95.9)	
	groups in heiGHt (Table 2). Subjects enrolled in fliGHt were primarily treatment-experienced (98%) and ranged from 1.2 to 17.4 years old	Withdrew from enliGHten*	3 (2.9)	1 (1.8)	4 (2.9)	
•	Upon entry into enliGHten, subjects from fliGHt were generally older and more advanced inTanner Stage	Completed enliGHten ^b	0	0	7 (5.0)	
	compared to those entering from heiGHt (Table 3)	Teconicator based on subjects exciled	and depend in emilipides			

Table 3. Demographics and Disease Characteristics at Start of enliGHte

		heiGH	heiGHt
		Lonapegaomatropin (n = 103)*	
	Age	Age	Age
	Mean (age)	Mean (age) 9.5 (2.7)	Mean (age) 9.5 (2.7) 9.5 (2.8)
	Min, max	Min, max 4.4, 14.1	Min, max 4.4, 14.1 4.2, 13.9
	Height SDS, Mean (SD)		
	Average IGF-1 SDS, mean (SD)		
-2.6 (1.3) -1.1 (1.0) Tanner Stage			
	Stage I, n (%)	Stage I, n (%) 92 (89.3)	Stage I, n (%) 92 (89.3) 45 (81.8)
	Stage II, n (%)	Stage II, n (%) 11 (10.7)	Stage II, n (%) 11 (10.7) 8 (14.5)
	Stage III, n (%)	Stage III, n (%) 0	Stage III, n (%) 0 2 (3.6)
	Stage IV, n (%)	Stage IV, n (%) 0	Stage IV, n (%) 0 0
	Stage V, n (%)	Stage V, n (%) 0	Stage V, n (%) 0 0
	Mean (age) Min, max Height SDS, Mean (SD) Average (GF1 SDS, mean (SD) Tanner Stage Stage I, n (%) Stage II, n (%) Stage II, n (%) Stage II, n (%)	Mann (age) Display Age 5.5 (2.7) Mann (age) 9.5 (2.7) Mann (age) 4.4, 14.1 Haigh 505, Marsh (30) 10.02, 17 Arrange 1671 505, marsh (30) 0.6 (0.3) Tanner Stage 11.102 Stage 1, n (%) 9.2 (0.9, 11 Stage 1, n (%) 0 Stage 1, n (%) 0	Idenge autors Descent autors Age 55 (27) 55 (28) Mon (age) 55 (27) 55 (28) Mon (age) 55 (27) 55 (28) Min max 45 (14) 42, 130 Height SDS. Mexicol 15 (02) 21 (08) Arrange (64) 55 (26) 65 (02) 201 (01) Tomore Stage 55 (28) 65 (02) 201 (01) Stage (1, 15) 10 (02) 21 (08) 56 (03) Stage (1, 15) 10 (02) 21 (08) 56 (03) Stage (1, 15) 0 (02) 21 (08) 56 (03) Stage (1, 15) 0 (02) 21 (03) 56 (03)

Height and Pharmacodynamic

- · For heiGHt subjects, AHV inc iects initially treated with lonapegasomat
- In heiGHt, average IGF-1 SD pared with daily somatropin-treated subjects
- stable without further increase; for heiGHt subjects who switched from daily somatropin to lonapegso

Figure 4. Average IGF-1 SDS over 104 Weeks for



· fliGHt subjects continued to approach their average parental height, with height SDS improving from -1.42 at fliGHt baseline to -0.69 at Week 78 (Figure 5); LS mean (SE) AHV at Week 78 was 8.3 cm/year and was consistent with clinical expectations given the characteristics of the enrolled subjects⁴

RESULTS

· As of this data cut, a total of 5 subjects have met or exceeded their target height (Table 4). All 5 subjects were from the fliGHt trial. Prior exposure to somatropin varied (range 0.6-2.4 years) but all received lonapegsomatropin for approximately 2 years (range 1.7-2.0 years). All 5 exceeded their average parental height SDS.

· For fliGHt subjects, observed mean (SD) average IGF-1 SDS increased from 0.85 (1.3) at fliGHt baseline to 1.62 (1.3) at Week 26 and 1.81 (1.1) at Week 78 (not shown)

Figure 5. Sustained Improvement in Height SDS (fliGHt trial)



Table 4. Subjects Who Met or Exceeded Target Height

Subject age (male)	Prior somatropin (years)	Duration on Ionapegao- matropin (years)	Height SDS at Baseline'	Last Height SDS	Avg Parental Height SDS*	Height at Baseline (cm) ⁷	Last Height (cm)	Target Height" (cm)	Last AHV* (cm/yr)
14.9	0.6	2.0	-1.6	-0.04	-0.37	156.5	174.9	174.0	8.8
15.4	2.1	1.8	-0.33	0.26	-0.97	169.2	177.3	170.3	2.8
15.6	0.5	1.8	-0.67	0.61	-0.83	167.2	180.0	171.0	5.8
14.4	2.4	1.7	-0.89	0.08	-0.54	159.6	173.9	172.9	9.3
15.6	1.0	2.0	-1.64	-0.23	-0.56	159.2	174.2	172.9	5.4

Safety Outcome

· The adverse event (AE) profile of lonapegsomatropin was consistent

- generally mild and no serious TEAEs were considered to be related to study drug (Table 5) In enliGHten, the most common TEAEs were upper respiratory tract infection (21,1%), nasopharyngitis (11,1%), cough (8,7%), and pyrexia (8.4%); these are consistent with other clinical trials evaluating daily somatropin in children with GHD⁶
- A low titer non-neutralizing anti-drug binding antibodies was detected in <10% of lonapegsomatropin-treated subjects (not shown)

across the phase 3 studies; treatment emergent AEs (TEAEs) were

- Hemoglobin A1c, cortisol, and free thyroxine were stable and generally remained within the normal range throughout the trials (not shown)
- Over time, treatment with lonapedsomatropin was associated with mean BMI SDS that stabilized towards 0 (Figure 6)

 Between the lonanegsomatropin and Genotropin groups, there was a similar change in bone age over 104 weeks (Table 6). The bone age/ chronological age ratios at Week 52 and Week 104 remained less than 1, epresenting normal skeletal maturation. Overall, this suggests that the longer-term effects of lonapegsomatropin (up to 104 weeks) did not occur at the expense of accelerated skeletal maturation.

Table 5. Summary of Adverse Events Across All Trials

2 (1.9)

0

Table 6. Similar Change in Bone Age Over 104 Weeks*

0.7 (0.2)

0.7 (0.1)

0.8 (0.1)

25(13)

23(14)

21(15)

Figure 6. BMI SDS Across All Trials

Bone age/chronological age ratio

Delay in bone age (years

Key Features

Room temperature storage

hGH (31G, 4mm)

· Simple operation

· Small needle, comparable to daily

· No waste due to empty-all design

Single low-volume (<0.60mL)

injection for patients <60kg

· Device lifespan at least 4 years

· Bluetooth® connectivity planned

connectivity platform underwa

for automatic data capture

Development of integrated

Enables flexible titration

105

104

98

105

104

98

Figure 7. TransCon hGH Auto-Injector Device

Baseline

Week 52

Week 104

Receline

Week 52

Week 104

1 (1.8)

Category, n (%)

Treatment-

Events (TEAEs)

TEAEs Related to

Study Drug

Study Drug

TEAEs Leadin

Study Drug

Study Drug

TEAEs Leading t

Serious Advers

heiGHt Trial (52 weeks)		fliGHt Trial (26 weeks)	enliGHten Trial (up to 52 weeks)
Lonapegaomatropin (n = 105)	Daily somatropin (n = 56)	Lonapegaomatropin (n = 146)	Lonapegaomatropin (N = 298)
81 (77)	39 (70)	83 (57)	195 (65.4)
12 (11)	10 (18)	6 (4.1)	13 (4.4)
1 (1.0)	1 (1.8)	1 (0.7)	10 (3.4)
0	0	0	0

2 (1.4)

0

0.7 (0.1)

0.8 (0.1)

2.3 (1.1)

-0.2 (0.9) 2.1 (1.1) -0.2 (0.7)

-0.4 (1.0) 2.0 (1.2) -0.4 (1.0)

0.8 (0.1) 0.1 (0.1)

0.1 (0.1)

0.1 (0.1)

0.1 (0.1)

() Mixing

5 (1.7)

Switching to the TransCon hGH Auto-Injecto

- Once available, subjects in enliGHten at select sites in the US were switched from vial and syringe to the TransCon hGH Auto-Injector; currently, 160 subjects are using the GH Auto-Injector in enliGHten (Figure 7)
- As indicated by the Device Usability Questionnaire, the TransCon hGH Auto-Injector was generally found to be comfortable, easy-to-use, and safe (Figure 8)
- Overall, fewer local tolerability reactions were reported with the TransCon hGH Auto-Injector compared with syringe/needle (Figure 9)

Figure 8. Device Usability Questionnaire (DUQ) Ratings at Week 6 After TransCon hGH Auto-Injector Initiation (n = 139)



Figure 9. Local Tolerability from Subject Diary



	switched to the TransCon hGH Aut			
	d any changes at the injection site,		bruising or itching" is a	depicted. Overall, fewer
injections given by the TransCo	in hGH Auto-Injector were rated as	such.		

CONCLUSIONS

- · Across the broad population of the phase 3 program, subjects treated with lonapegsomatropin for up to 2 years continued to grow well, with a safety profile comparable to daily growth hormone, including a similar AE profile, stable BMI, stable laboratory parameters, and low immunogenicity
- · Among subjects who switched from daily somatropin to lonapegsomatropin, a lower-than-expected attenuation in 2nd year AHV suggested an improved treatment effect of lonapegsomatropin relative to the previous daily
- · As of this data cut, 5 subjects have met or exceeded their target height and have also exceeded their average parental height SDS
- High retention rates were observed across the phase 3 program, with >98% of subjects continuing from heiGHt and fliGHt into enliGHte
- · The TransCon hGH Auto-Injector was found to be well-tolerated and easy-touse by caregivers and subjects

ENDO June 2020. 5. Bakker B et al.

cendis, Ascendis Pharma, Ascendis Pharma logo, the company logo and TransCon are trademarks

Outcomes
creased from an untreated baseline of 3.93 cm/year in favor of subje ropin (Figure 3)
S values were higher for lonapegsomatropin-treated subjects comp , paralleling the observed improved growth outcomes (Figure 4)

· Beyond 52 weeks, average IGF-1 SDS for heiGHt subjects who started on lonapegsomatropin generally remained an initial increase in average IGF-1 SDS with subsequent stabilization was observed (Figure 4)



- Figure 3. AHV over 104 Weeks for heiGHt subjects Comparisons between the two heiGHt treatment groups allowed for the evaluation of safety and
- efficacy outcomes as they had similar baseline demographics and comparable treatment histories IGE-1 was obtained on post-dose Day 5 (+1) in fliGHt and enliGHten: in heiGHt, average IGE-1 for lonapegsomatropin was calculated based on a

population pharmacodynamic model SAFETY

 Safety was evaluated throughout the trial periods and is summarized by trial

STATISTICAL ANALYSIS

· A by-visit ANCOVA model was used to analyze numeric endpoints

21 (37.5)	52 (35.6)		Stage IV, n (%)	0	0	17 (
35 (62.5) 91 (62.3)			Stage V, n (%)	0	0	3 (
rans to SIGHT; subjects had been previously treated			baseline sample size subject to minor insignili oliculated using a pharmacodynamic model	cart dange from parameters to p	asmeters	

68 (64.8)