Effects of Solriamfetol on Cognition in Obstructive Sleep Apnea With **Excessive Daytime Sleepiness and** Impaired Cognition in the SHARP **Clinical Trial**

Hans P.A. Van Dongen, PhD¹; Eileen B. Leary, PhD, RPSGT²; Graham M.L. Eglit, PhD²; Sunayana B. Banerjee, PhD²; Christopher Drake, PhD³; Richard Bogan, MD, FCCP⁴; Judith Jaeger, PhD, MPA⁵; Herriot Tabuteau, MD²

¹Department of Translational Medicine and Physiology & Sleep and Performance Research Center, Washington State University, Spokane, WA, USA; ²Axsome Therapeutics, New York, NY, USA; ³Henry Ford Health System, Detroit, MI, USA; ⁴SleepMed, Inc., Columbia, SC, USA; ⁵CognitionMetrics, Stamford, CT, USA

Key Questions

- What are the effects of solriamfetol treatment on individual cognitive complaints related to concentration, memory and thinking skills as measured by the British Columbia Cognitive Complaints Inventory (BC-CCI)?
- What are the effects of solriamfetol treatment on individual functional items of the BC-CCI that assess social functioning, vocational functioning, and management of personal relationships?

Conclusions

- Consistent with previous reports showing improvement on objective cognitive measures, solriamfetol led to significant subjective improvements overall, and particularly in subjective cognitive domains that may be related to **memory, executive functioning,** and processing speed
- Solriamfetol has the potential to improve subjective cognitive functioning in participants with impaired cognition associated with USA and EDS

References

- . Gasa M, et al. J Sleep Res. 2013;22(4):389-97.
- 2. Pepin JL, et al. Eur Respir J. 2009;33(5):1062-7
- 3. Bonsignore MR, et al. Front Neurol. 2021;12:690008.
- 4. Zhou J, et al. *Sleep Med.* 2016;23:99-108.
- 5. Vasudev P, et al. Indian J Sleep Med. 2020;15(4):55-9.
- 6. Gursahani H, et al. *Sleep.* 2022;45(suppl 1):A329.
- 7. Sunosi[®] (solriamfetol) tablets [prescribing information]. New York, NY: Axsome Therapeutics, Inc; 2022. 8. Sunosi[®] (solriamfetol) tablets Summary of Product Characteristics. København S, Denmark: Atnahs Pharma
- Netherlands B. V.; 2023.
- 9. Sunosi [product monograph including patient medical information]. Malta: Axsome Malta Ltd.; 2022.
- 10. lverson GL, Lam RW. Ann Clin Psychiatry. 2013;25(2):135-40.

Acknowledgments

This study was supported by Axsome Therapeutics, Inc. and Jazz Pharmaceuticals. The development of this presentation was supported by Axsome Therapeutics, Inc. Under the direction of the authors, Jacob Huffman, PhD, of Peloton Advantage, LLC, an OPEN Health company, provided medical writing and editorial support for this poster, which was funded by Axsome Therapeutics, Inc. The authors thank the participants who contributed data, personnel who collected data, and consultants who contributed to the design of the SHARP Study.

Disclosures

H.P.A. Van Dongen serves as a paid consultant to Jazz Pharmaceuticals.

E.B. Leary was employed by Jazz Pharmaceuticals during the time the study was conducted and is a current employee of Axsome Therapeutics. C. Drake serves as a consultant to Axsome, Harmony, Takeda, Procter & Gamble, Apnimed, Zevra – Research; Harmony, Idorsia, Reunion – Speaker; Procter & Gamble, and Zevra.

R. Bogan: serves as a consultant to Axsome Therapeutics, Avadel, Harmony, Jazz Pharmaceuticals, and Takeda and is on the speakers bureau for Axsome Therapeutics, Harmony, Idorsia, and Jazz Pharmaceuticals.

J. Jaeger is an employee of Cognition Metrics, LLC; Cognition Metrics received research support from Jazz Pharmaceuticals and Axsome [herapeutics.

G. Eglit and H. Tabuteau are current employees of Axsome Therapeutics.



an QR code or access

ps://www.axsomecongresshub.com/NEI2024 o view or download a PDF of this poster or access dditional information and other Axsome Therapeutics presentations at NEI 2024.



Neuroscience Education Institute (NEI) Congress November 7-10, 2024, Colorado Springs, CO

Introduction

- cognitive domains⁴⁻⁵
- Solriamfetol (Sunosi[®]) is a dopamine and norepinephrine reuptake inhibitor with agonistic properties at trace amine-associated receptor 1 (TAAR1) and serotonin 1A receptors⁶⁻⁷
- Solriamfetol is approved in the United States, Canada, and select European countries to treat EDS associated with OSA (37.5–150 mg)

SHARP Trial (NCT04789174)

- impaired cognition

Key Findings

Table 1. Baseline Demographics and Clinical Characteristics

Age, mean (SD), years Sex (female), n (%) Race, n (%) White Black/African Americar Asian Unknown Body mass index, mean (Digit Symbol Substitution BC-CCI, mean (SD) Patient Global Impression Epworth Sleepiness Scale Positive airway pressure Adherent use (≥4 h/ni Hours of use (among al Table 2. Baseline Scores on Individual BC-CCI Items

Cognitive complaint items	
complaint items	
Functional items	

Excessive daytime sleepiness (EDS) is common in patients with obstructive sleep apnea (OSA), and can persist in up to 28% of patients despite use of primary airway therapy¹⁻³

Patients with EDS associated with OSA can have deficits in several

Objective: to assess whether solriamfetol improves cognitive function in patients with EDS associated with OSA and extant

This post hoc analysis evaluated the effects of solriamfetol on individual cognitive complaints and functional items on the British **Columbia Cognitive Complaints Inventory (BC-CCI)**

Methods & Study Design





• Of 173 participants screened, 59 were enrolled and had baseline data, 58 had data available for efficacy analyses, and 57 completed the study

Baseline characteristics, including baseline total BC-CCI scores, were generally similar between groups – Baseline scores on individual BC-CCI items were generally similar between groups

■ Among participants using positive airway pressure, average use was ≥6 hours per night

	Solriamfetol/ placebo (n=30)	Placebo/ solriamfetol (n=29)	Overall (N=59)
	52.5 (10.5)	51.9 (11.1)	52.2 (10.7)
	10 (33.3)	11 (37.9)	21 (35.6)
	24 (80.0)	19 (65.5)	43 (72.9)
an	4 (13.3)	8 (27.6)	12 (20.3)
	1 (3.3)	2 (6.9)	3 (5.1)
	1 (3.3)	0	1 (1.7)
(SD), kg/m ²	32.8 (4.7)	31.6 (4.0)	32.2 (4.4)
n Test, age-corrected, mean (SD)	6.6 (1.3)	6.9 (0.8)	6.8 (1.1)
	11.4 (2.5)	11.4 (2.5)	11.4 (2.5)
n of Severity (cognitive function), mean (SD)	2.2 (0.8)	2.3 (0.7)	2.3 (0.7)
e total score, mean (SD)	14.8 (2.8)	14.3 (2.7)	14.6 (2.8)
use, n (%)	22 (73.3)	20 (69.0)	42 (71.2)
ight for 70% of nights), n (%)	18 (60.0)	16 (55.2)	34 (57.6)
all users), mean (SD)	6.0 (2.4)	6.6 (2.7)	6.3 (2.5)

Baseline scores on individual BC-CCI items were generally similar for participants randomized to solriamfetol/placebo versus placebo/solriamfetol

Mean (SD)	Solriamfetol/ placebo (n=29)	Placebo/ solriamfetol (n=29)	Overall (N=58)
Forgetfulness/memory problems	1.93 (0.70)	2.00 (0.71)	1.97 (0.70)
Poor concentration	2.10 (0.86)	2.21 (0.68)	2.16 (0.77)
Trouble expressing thoughts	1.93 (0.80)	1.76 (0.74)	1.84 (0.77)
Trouble finding the right word	1.97 (0.82)	1.79 (0.56)	1.88 (0.70)
Slow thinking speed	1.93 (0.75)	1.93 (0.80)	1.93 (0.77)
Trouble figuring things out	1.62 (0.73)	1.76 (0.64)	1.69 (0.68)
Vocational functioning	1.97 (1.05)	1.83 (0.97)	1.90 (1.00)
Family/friends functioning	1.52 (1.02)	1.72 (1.16)	1.62 (1.09)
Social/recreational functioning	1.66 (1.11)	1.66 (1.04)	1.66 (1.07)

Figure 4. Primary Findings and Safety



Figure 5. Overall Improvement in Subjective Cognitive Function

function) after solriamfetol treatment compared with placebo



• Overall, BC-CCI scores showed greater reduction from baseline (ie, more improvement in subjective cognitive

mprovement in subjective cognitive function

Solriamfetol significantly improved subjective cognitive function compared with placebo

Least squares mean difference: -1.58 (95% CI: -2.53, -0.63) Cohen's *d*: 0.45

Figure 6. Cognitive Complaint Items

Cognitive Complaint Items		Difference From Placebo (95% Cl)	Cohen's d	1
Poor concentration –	<i>P</i> =0.007	-0.36 (-0.62, -0.10)	d= 0.37	Solriamfetol
Slow thinking speed –	<i>P</i> =0.009	-0.31 (-0.54, -0.08)	d= 0.36	significantly improved most
Trouble finding the right word –	<i>P</i> =0.042	-0.24 (-0.48, -0.01)	d= 0.28	individual cognitive complaint items
Trouble figuring things out –	<i>P</i> =0.030	-0.23 (-0.43, -0.02)	d= 0.30	compared with placebo
Forgetfulness/memory problems –	<i>P</i> =0.013	-0.22 (-0.40, -0.05)	d= 0.34	
Trouble expressing thoughts –	<i>P</i> =0.077	-0.21 (-0.44, 0.02)	<i>d</i> =0.24	
-0.	8 -0.6 -0.4 -0.2 0.4	0 0.2		
	Difference From Placebo)		

Figure 7. Functional Items



	3 Functional Items
ms with during	Participants were asked to answer questions about how the cognitive complaints impacted their ability to function in the last 7 days. Questions included:
	1. Symptoms made it difficult to do job
	 Symptoms made it difficult to have good relationships with family and friends
	3. Symptoms made it difficult to enjoy social activities, recreational activities, or hobbies
oblems scores	Answer options included: False, Not at all true Slightly true Mainly true Very true
	Complaints for the BC-CCI Total Scores

Classifications for Cognitive Complaints for the BC-CCI Total Scores Calculated as the sum of the 6 cognitive complaint responses

	• 9 to 14: "moderate" cognitive complaints
complaints	• 15 to 18: "severe" cognitive complaints