# Real-world Use of Solriamfetol for **Excessive Daytime Sleepiness in** Patients Reporting Anxiety or Depression in the Real-World **SURWEY Study**

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# **Key Question**

■ Is solriamfetol effective in treating excessive daytime sleepiness associated with narcolepsy or obstructive sleep apnea in patients with self-reported anxiety and/or depression?

# Conclusions

- These real-world data describe treatment outcomes of solriamfetol in patients with narcolepsy or OSA, both with and without self-reported anxiety/depression
- Reductions in EDS were substantial and comparable in patients with and without self-reported anxiety/depression
- Most patients and physicians reported improvements in EDS
- These findings are consistent with clinical trial results and suggest that solriamfetol is effective in managing EDS in patients with psychiatric comorbidities

## eferences

- . Fortuyn H, et al. *Gen Hosp Psychiatry*. 2010;32(1):49-56.
- . Sharafkhaneh A. et al. *Sleep*. 2005 28(11):1405-11.
- Kim JY, et al. JAMA Otolaryngol Head Neck Surg. 2019;145(11):1020-1026.
- 4. Garbarino S, et al. Behav Sleep Med. 2020; 18(1):35-57. 5. Alnefeesi Y, et al. Neurosci Biobehav Rev. 2021;131:192-210
- 6. Gursahani H, et al W. Sleep. 2022;45(suppl 1):A329
- 7. Sunosi® (solriamfetol) [Prescribing Information]. New York, NY. Axsome Therapeutics, Inc.

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8. Sunosi™ (solriamfetol) tablets Summary of Product Characteristics. Waterford, Ireland: TMC Pharma (EU) Limited; 2022.

G.M.L. Eglit is an employee of Axsome Therapeutics, Inc.

S. Floam is an employee of Axsome Therapeutics, Inc. and former employees of Jazz Pharmaceuticals.

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# Introduction

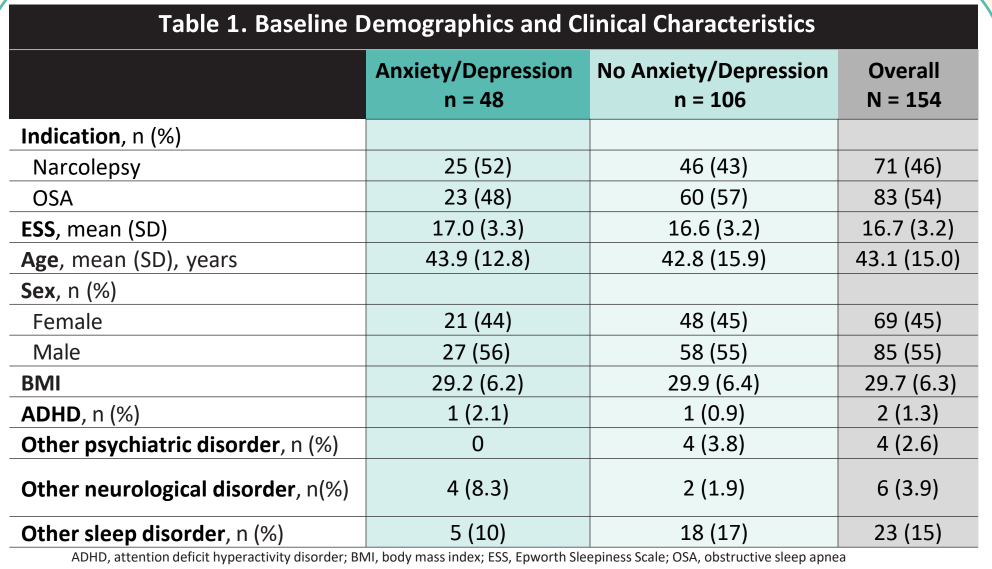
- Psychiatric comorbidities are prevalent in patients with excessive daytime sleepiness (EDS) from narcolepsy or obstructive sleep apnea (OSA)<sup>1,2</sup> - Depression and anxiety are particularly common in these patients, with prevalence rates of  $\geq$  30% each<sup>3,4</sup>
- Efficacy and safety data for wake-promoting agents in these populations are limited
- Solriamfetol (Sunosi®) is a dopamine-norepinephrine reuptake inhibitor with agonistic properties at the trace amine-associated receptor 1 and serotonin 1A receptor<sup>5,6</sup>; it is approved for use in adults in the United States, Canada and select countries in Europe for the treatment of EDS associated with narcolepsy or OSA<sup>7,8</sup>
- Clinical trials with solriamfetol have excluded patients with severe psychiatric comorbidities, and the prescribing information advises against its use in this population
- As a result, there are limited data available on the efficacy and safety of solriamfetol in these patients

# **Methods & Study Design**

- **SU**nosi **R**eal **W**orld **E**xperience Stud**Y** (SURWEY) was a retrospective chart review among physicians in Germany who have prescribed solriamfetol to patients with EDS associated with narcolepsy or OSA
- Eligible patients were ≥ 18 years of age, had a diagnosis of EDS and narcolepsy or OSA, had reached a stable maintenance dose of solriamfetol and completed ≥ 6 weeks of treatment; patients who received solriamfetol during a clinical trial or early access program were excluded
- The present analysis focused on data from 154 adult patients with narcolepsy or OSA, stratified by self-reported anxiety and/or depression Patients were classified as anxious and/or depressed based on their answer at baseline to a single yes/no question
- Data related to comorbidities, Epworth Sleepiness Scale (ESS) scores, patient-and physician-reported improvement in EDS, and adverse events were summarized descriptively

# **Key Findings**

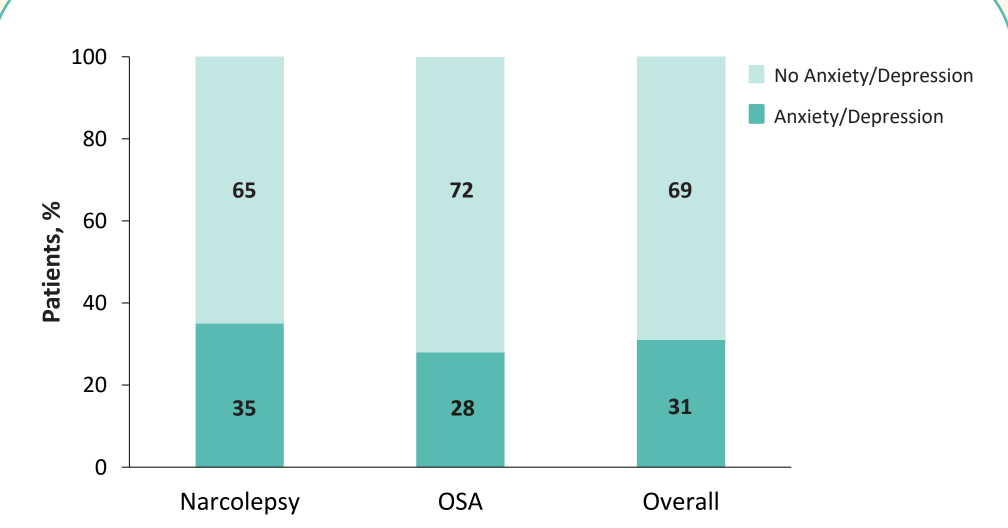
**Patient Populatio** 



Baseline demographics were similar between patients with and without self-reported anxiety and/or depression

## **Anxiety/Depression Incidence**

# Figure 1. Incidence of Anxiety/Depression in Patients With Narcolepsy or OSA



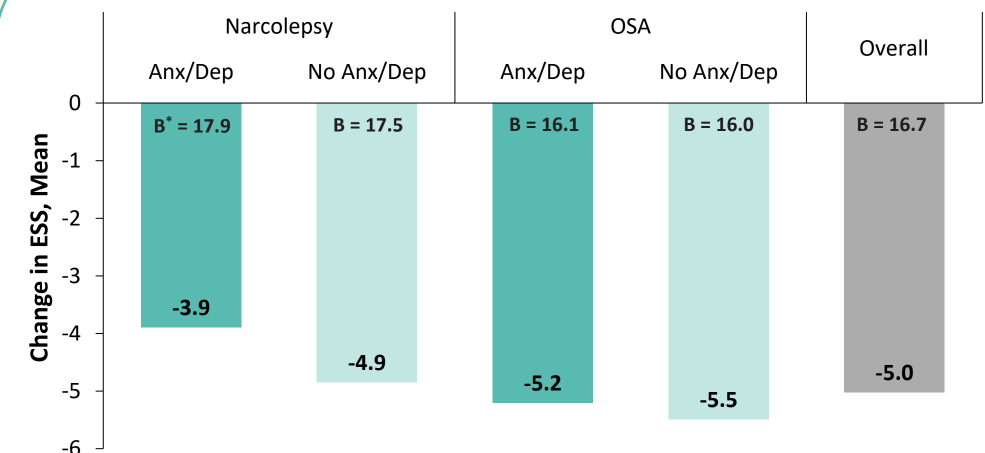
 Rates of anxiety/depression were similar between patients with narcolepsy (35.2%) and OSA (27.7%)

# **Efficacy**

 All efficacy results were pooled across dosages, and most patients took less than the maximum recommended dose of 150mg/day

**Average Improvements in EDS** 

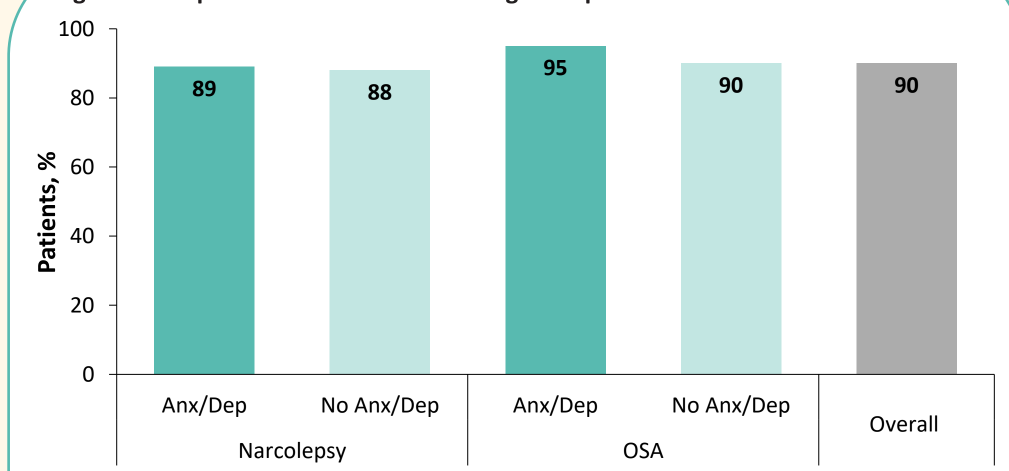
Figure 2. Reductions in ESS Scores for Patients With and Without Anxiety/Depression



 In patients with narcolepsy or OSA, those with anxiety/depression experienced comparable reductions in ESS to those without

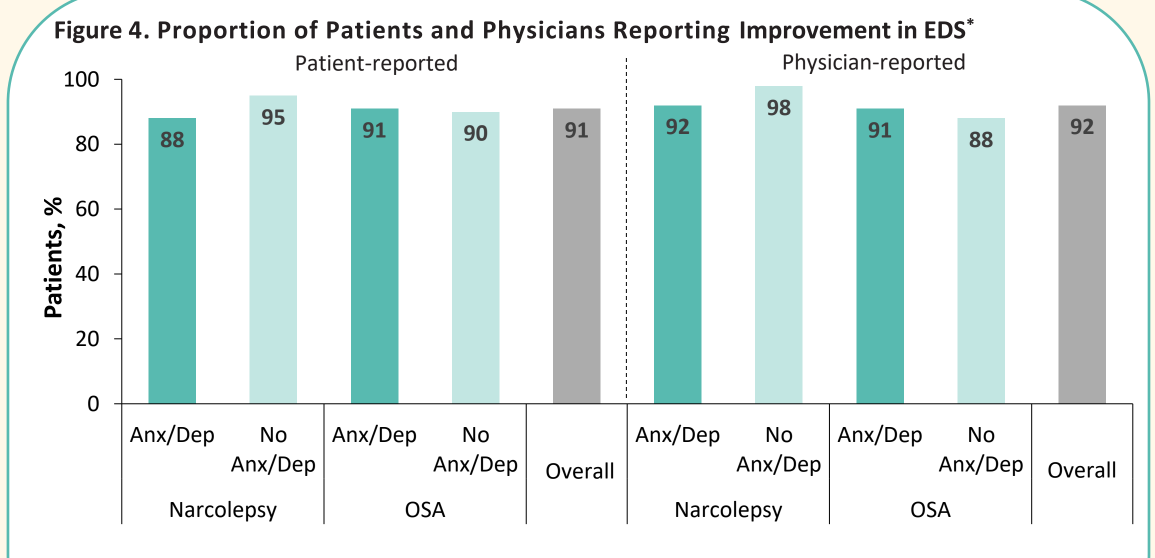
# Clinically Meaningful Improvement

Figure 3. Proportion of Patients Achieving a ≥ 2-point Reduction in ESS Score



• In patients with narcolepsy or OSA, ≥ 88% experienced clinically meaningful improvement in EDS, achieving a reduction of ≥ 2 points in ESS score, regardless of anxiety/depression status

# Patient/Physician-Reported Improvemen



In patients with narcolepsy or OSA, ≥ 88% reported experiencing improvement in EDS, regardless of anxiety/depression status, consistent with physician reports

\*Patients or physicians rated EDS "slightly improved" or "strongly improved"

Safety

## **Table 2. Adverse Events (≥3% Overall) OSA** Overall Narcolepsy No anxiety/ Anxiety/ Anxiety/ No anxiety/ depression depression depression depression n = 46n = 23 n = 60 N = 154Adverse event, n (%) n = 25 2 (8.3) 3 (13.0) 4 (8.9) 4 (6.8) 13 (8.6) Headache **Decreased** 1 (4.2) 10 (6.6) 3 (6.7) 3 (13.0) 3 (5.1) appetite 2 (8.3) 2 (8.7) 9 (6.0) 2 (4.4) 3 (5.1) Insomnia 3 (12.5) 2 (8.7) 2 (3.4) 7 (4.6) Irritability 3 (12.5) 6 (4.0) 3 (5.1) Other 1 (4.2) 2 (3.4) 5 (3.3) 1 (2.2) 1 (4.3) Dizziness **Feeling jittery** 1 (4.2) 1 (4.3) 3 (5.1) 5 (3.3)

- The most common adverse events overall were headache, decreased appetite, and insomnia (**Table 2**)
- Adverse events were more common overall in patients reporting anxiety/depression (Table 2)