

FOR ADULTS WITH EXCESSIVE DAYTIME SLEEPINESS (EDS) DUE TO OBSTRUCTIVE SLEEP APNEA (OSA)¹



THE SUNOSI CHALLENGE

Help your patients identify the treatment option that is best for them*

Identify



Identify which of your commercially insured adult patients diagnosed with OSA continue to have excessive daytime sleepiness despite adherence with airway therapy, such as CPAP or oral appliance

Initiate



Initiate a **SUNOSI trial** for new and existing patients with excessive daytime sleepiness due to OSA[†]

Talk



Talk with your patients about their experience. If they decide to continue with SUNOSI, you can prescribe up to **90 days for as little as \$9** with the savings offer[‡]

Initiating the SUNOSI trial:

Provide SUNOSI samples or a free trial offer[†] at no cost to your patients

CPAP=continuous positive airway pressure.

*The content herein is not an attempt to practice medicine or provide specific medical advice, and it is not intended to make a diagnosis or replace or overrule a qualified healthcare provider's judgment. The final decision about whether to start, continue, or stop medical treatment rests with the healthcare provider, based on their professional assessment of the patient's condition, medical history, and the best course of action, considering all relevant factors.

[†]A valid prescription is required. Patients may receive a lifetime maximum of 30 tablets of SUNOSI (75 mg and 150 mg only) through this free trial offer. See full Terms and Conditions at <https://www.sunosihcp.com/savings-card-and-coverage>

[‡]Eligible patients will pay as little as \$9 with a valid prescription for an FDA-approved indication; monthly, annual, and/or per-claim maximum program benefits may apply and vary depending on the patients' specific terms of their prescription drug plan and to ensure that the funds are used for the benefit of the patient, based on factors determined by Axxome. [See full Terms and Conditions.](#) Some insurance plans may require a step therapy of modafinil or armodafinil.

INDICATION AND IMPORTANT SAFETY INFORMATION FOR SUNOSI (solriamfetol)

INDICATION

SUNOSI is indicated to improve wakefulness in adults with excessive daytime sleepiness (EDS) associated with obstructive sleep apnea (OSA).

LIMITATIONS OF USE

SUNOSI is not indicated to treat the underlying obstruction in OSA. Ensure that the underlying airway obstruction is treated (e.g., with continuous positive airway pressure (CPAP)) for at least one month prior to initiating SUNOSI. SUNOSI is not a substitute for these modalities, and the treatment of the underlying airway obstruction should be continued.

CONTRAINDICATIONS

SUNOSI is contraindicated in patients receiving concomitant treatment with monoamine oxidase inhibitors (MAOIs), or within 14 days following discontinuation of an MAOI, because of the risk of hypertensive reaction.

Please see next page for additional Important Safety Information and full [Prescribing Information.](#)



Which of your appropriate patients are ready for the Challenge?



IMPORTANT SAFETY INFORMATION (CONT'D)

WARNINGS AND PRECAUTIONS

Blood Pressure and Heart Rate Increases

SUNOSI increases systolic blood pressure, diastolic blood pressure, and heart rate in a dose-dependent fashion.

Epidemiological data show that chronic elevations in blood pressure increase the risk of major adverse cardiovascular events (MACE), including stroke, heart attack, and cardiovascular death. The magnitude of the increase in absolute risk is dependent on the increase in blood pressure and the underlying risk of MACE in the population being treated. Many patients with narcolepsy and OSA have multiple risk factors for MACE, including hypertension, diabetes, hyperlipidemia, and high body mass index (BMI).

Assess blood pressure and control hypertension before initiating treatment with SUNOSI. Monitor blood pressure regularly during treatment and treat new-onset hypertension and exacerbations of pre-existing hypertension. Exercise caution when treating patients at higher risk of MACE, particularly patients with known cardiovascular and cerebrovascular disease, pre-existing hypertension, and patients with advanced age. Use caution with other drugs that increase blood pressure and heart rate.

Periodically reassess the need for continued treatment with SUNOSI. If a patient experiences increases in blood pressure or heart rate that cannot be managed with dose reduction of SUNOSI or other appropriate medical intervention, consider discontinuation of SUNOSI.

Patients with moderate or severe renal impairment could be at a higher risk of increases in blood pressure and heart rate because of the prolonged half-life of SUNOSI.

Psychiatric Symptoms

Psychiatric adverse reactions have been observed in clinical trials with SUNOSI, including anxiety, insomnia, and irritability.

Exercise caution when treating patients with SUNOSI who have a history of psychosis or bipolar disorders, as SUNOSI has not been evaluated in these patients.

Patients with moderate or severe renal impairment may be at a higher risk of psychiatric symptoms because of the prolonged half-life of SUNOSI.

Observe SUNOSI patients for the possible emergence or exacerbation of psychiatric symptoms. Consider dose reduction or discontinuation of SUNOSI if psychiatric symptoms develop.

MOST COMMON ADVERSE REACTIONS

The most common adverse reactions (incidence $\geq 5\%$) reported more frequently with the use of SUNOSI than placebo in either narcolepsy or OSA were headache, nausea, decreased appetite, anxiety, and insomnia.

SUN HCP aISI 05/2022

Please see full [Prescribing Information](#).

Reference:

1. SUNOSI (solriamfetol) [prescribing information]. New York, NY: Axsome Therapeutics, Inc.

axsome

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SUNOSI
(solriamfetol) ^{IV}
75, 150 mg tablets