

YOU ARE CORDIALLY INVITED TO ATTEND A PEER-TO-PEER PRESENTATION

Spotlight on XCOPRI® (cenobamate tablets) CV for the Treatment of Partial-Onset Seizures in Adult Patients

XCOPRI®
(cenobamate tablets) CV
12.5 • 25 • 50 • 100 • 150 • 200 mg

PROGRAM DETAILS

Wednesday, April 14, 2021

6:30 PM CT

Virtual

Virtual

During this program, our presenter will discuss the following:

- The XCOPRI clinical development program
- Efficacy and safety of XCOPRI
- Considerations when initiating XCOPRI

Should you accept a meal while attending this speaker program, you are accepting the disclosure of the cost of the meal. Please note that meal cost limits vary by state, and it is the responsibility of individual attendees to adhere to specific state requirements.

PRESENTED BY

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REGISTER TODAY!

Please register by going to: <https://scimedregister.com>

Please reference program code: 2EWSEM

This is considered an SKLSI promotional program and branded materials will be provided, therefore all FDA promotional rules apply. XCOPRI is indicated for the treatment of partial-onset seizures in adult patients. For any questions regarding XCOPRI, please refer to the full [Prescribing Information](#) and Important Safety Information.

To unsubscribe or opt out of receiving future messages regarding this event, please email SKLifeSciences@scimedny.com.

SK Life Science is required to disclose all items of value provided to healthcare providers and to disclose these amounts publicly. By attending this speaker program, you are accepting the disclosure of the cost of the meal. Please note that meal cost limits vary by state, and it is the responsibility of individual attendees to adhere to specific state requirements.

IMPORTANT SAFETY INFORMATION and INDICATION FOR XCOPRI (cenobamate tablets) CV

CONTRAINDICATIONS

XCOPRI is contraindicated in any patients with known hypersensitivity to the compound or any of the components of the drug product.

XCOPRI is contraindicated in patients with Familial Short QT syndrome.

Please see additional Important Safety Information on next page.

IMPORTANT SAFETY INFORMATION and INDICATION FOR XCOPRI (cenobamate tablets) CV (cont.)

WARNINGS AND PRECAUTIONS

Drug Reaction with Eosinophilia and Systemic Symptoms (DRESS): Also known as Multiorgan hypersensitivity, has been reported in patients taking antiepileptic drugs, including XCOPRI. DRESS has been reported, including one fatality, when XCOPRI is titrated rapidly (weekly or faster titration). No cases of DRESS were reported in an open-label safety study of 1339 partial-onset seizure patients when XCOPRI was initiated at 12.5 mg/day and titrated every two weeks. This finding does not establish that the risk of DRESS is prevented by a slower titration; however, XCOPRI should be initiated at 12.5 mg once daily and titrated every two weeks. DRESS typically, although not exclusively, presents with fever, rash, and/or lymphadenopathy, in association with other organ system involvement. Eosinophilia is often present. If such signs or symptoms are present, the patient should be evaluated immediately. XCOPRI should be discontinued immediately and not restarted if an alternative etiology for the signs or symptoms cannot be established.

QT Shortening: XCOPRI can cause shortening of the QT interval. Caution should be used when administering XCOPRI and other drugs that shorten the QT interval as there may be a synergistic effect on the QT interval that would increase the QT shortening risk.

Suicidal Behavior and Ideation: Antiepileptic drugs (AEDs), including XCOPRI, increase the risk of suicidal thoughts or behavior in patients taking these drugs for any indication. Patients treated with any AED for any indication should be monitored for the emergence or worsening of depression, suicidal thoughts or behavior, and/or any unusual changes in mood or behavior. Advise patients, their caregivers, and/or families to be alert for these behavioral changes and report them immediately to a healthcare provider.

Neurological Adverse Reactions: XCOPRI causes dose-dependent increases in the neurologic adverse reactions including, dizziness, diplopia, disturbance in gait and coordination, somnolence, and fatigue.

Prescribers should advise patients against engaging in hazardous activities requiring mental alertness, such as operating motor vehicles or dangerous machinery, until the effect of XCOPRI is known.

Withdrawal of AEDs: As with all antiepileptic drugs, XCOPRI should generally be withdrawn gradually because of the risk of increased seizure frequency and status epilepticus. But if withdrawal is needed because of a serious adverse event, rapid discontinuation can be considered.

MOST COMMON ADVERSE REACTIONS

In adult adjunctive therapy placebo-controlled clinical studies, the most common adverse reactions that occurred in XCOPRI-treated patients (incidence at least 10% and greater than placebo) were somnolence, dizziness, fatigue, diplopia, headache.

DOSING CONSIDERATIONS

Dosage adjustment of XCOPRI or other concomitant medications may be necessary.

- Consider gradually reducing phenytoin dosages by up to 50% during initial titration.
- Consider reducing dosages of phenobarbital and clobazam as needed when used concomitantly with XCOPRI. When XCOPRI and carbamazepine or lamotrigine are taken concomitantly, consider increasing dosages as needed of carbamazepine or lamotrigine.
- Consider increasing dosages as needed of drugs which are CYP2B6 and CYP3A substrates and decreasing dosages as needed of drugs which are CYP2C19 substrates.
- Effectiveness of hormonal oral contraceptives may be reduced when administered concomitantly with XCOPRI. Women should use additional or alternative non-hormonal birth control.

Dosage reduction of XCOPRI may be considered in patients with mild to moderate and severe renal impairment. XCOPRI use is not recommended in end-stage renal disease.

The maximum recommended daily dose is 200 mg for patients with mild or moderate hepatic impairment. XCOPRI use is not recommended in patients with severe hepatic impairment.

DRUG ABUSE

XCOPRI is a Schedule V controlled substance.

INDICATION

XCOPRI is indicated for the treatment of partial-onset seizures in adult patients.

Please see full [Prescribing Information](#).