Please complete and sign this form. Fax completed form to 1-866-217-8034.

THE FOLLOWING SECTION IS TO BE FILLED OUT BY YOUR DOCTOR’S OFFICE

Prescriber Information
Prescriber Name: ___________________________ Neurologist/Epileptologist Other: _______________ NPI # _________ DEA # _________
Facility Name/Address: ________________________ City: _______________ State: ___________ ZIP: ___________
Office Contact Name: ________________________ Email: ________________________ Phone: (       ) _______ Fax: (       ) _______

Clinical Information for Insurance Prior Authorizations— Please include a copy of patient’s clinical notes, if available.

Diagnosis
Diagnosis code(s):

Most Recent Antiepileptic Drug Treatment
Approximate Start and End Dates of Most Recent Treatment:
Surgical History:
Drug Allergies:

Prescription (Circle one option for each section)

VALTOCO Dosage Strength: 5 mg 10 mg 15 mg 20 mg

Instructions: ____________________________

Prescriber Authorization (Required)
I authorize the designated pharmacy to act as an agent to initiate and execute the insurance prior authorization process, if necessary, for this prescription and any future fills of the same prescription for the patient listed above. I understand that I can revoke this designation at any time by providing written notice.

Prescriber’s Signature: ____________________________ Date ___________ / _______

Please see accompanying full Prescribing Information, including Boxed Warning, and full Important Safety Information on the back.
Indication
VALTOCO® (diazepam nasal spray) is indicated for the acute treatment of intermittent, stereotypic episodes of frequent seizure activity (ie, seizure clusters, acute repetitive seizures) that are distinct from a patient's usual seizure pattern in patients with epilepsy 6 years of age and older.

IMPORTANT SAFETY INFORMATION

RISK FROM CONCOMITANT USE WITH OPIOIDS
Concomitant use of benzodiazepines and opioids may result in profound sedation, respiratory depression, coma, and death.
- Reserve concomitant prescribing of these drugs for use in patients for whom alternative treatment options are inadequate
- Limit dosages and durations to the minimum required
- Follow patients for signs and symptoms of respiratory depression and sedation

Contraindications: VALTOCO is contraindicated in patients with:
- Known hypersensitivity to diazepam
- Acute narrow-angle glaucoma

Central Nervous System (CNS) Depression
Benzodiazepines, including VALTOCO, may produce CNS depression. Caution patients against engaging in hazardous activities requiring mental alertness, such as operating machinery, driving a motor vehicle, or riding a bicycle, until the effects of the drug, such as drowsiness, have subsided, and as their medical condition permits.

The potential for a synergistic CNS-depressant effect when VALTOCO is used with alcohol or other CNS depressants must be considered, and appropriate recommendations made to the patient and/or care partner.

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

Glaucoma
Benzodiazepines, including VALTOCO, can increase intraocular pressure in patients with glaucoma. VALTOCO may only be used in patients with open-angle glaucoma only if they are receiving appropriate therapy. VALTOCO is contraindicated in patients with narrow-angle glaucoma.

Risk of Serious Adverse Reactions in Infants due to Benzyl Alcohol Preservative
VALTOCO is not approved for use in neonates or infants. Serious and fatal adverse reactions, including “gaspig syndrome,” can occur in neonates and low-birth-weight infants treated with benzyl alcohol-preserved drugs, including VALTOCO. The “gaspig syndrome” is characterized by central nervous system depression, metabolic acidosis, and gasping respirations. The minimum amount of benzyl alcohol at which serious adverse reactions may occur is not known.

Adverse Reactions
The most common adverse reactions (at least 4%) were somnolence, headache, and nasal discomfort.

Diazepam, the active ingredient in VALTOCO, is a Schedule IV controlled substance.

To report SUSPECTED ADVERSE REACTIONS, contact Neurelis, Inc. at 1-866-696-3873 or FDA at 1-800-FDA-1088 (www.fda.gov/medwatch).

Please read full Prescribing Information, including Boxed Warning, for additional important safety information.