

Vutrisiran

(Amvuttra[®]) J0225 (1 mg)

Covered with prior authorization

Initial Requests

Vutrisiran (Amvuttra[®]) may be authorized when the following criteria are met:

Individual is 18 years of age or older; **AND**

Individual has a diagnosis of hereditary transthyretin (hATTR) amyloidosis or familial amyloid polyneuropathy (FAP); **AND**

Documentation is provided that individual has a TTR mutation confirmed by genotyping **AND**

Documentation is provided that individual has associated mild to moderate polyneuropathy. (excluding diabetic)

Continuation of therapy

Vutrisiran (Amvuttra[®]) may be authorized when the following criteria are met:

Documentation is provided that there is confirmation of clinically significant improvement or stabilization in clinical signs and symptoms of disease (including but not limited to improved ambulation, improvement in neurologic symptom burden, improvement in activities of daily living.)

Exclusion criteria:

Requests for Vutrisiran (Amvuttra[®]) may not be approved for the following:

- Individual has a history of liver transplantation
- Individual is pregnant
- Individual has severe renal impairment or end-stage renal disease
- Individual has moderate or severe hepatic impairment
- Individual has New York Heart Association (NYHA) class III or IV heart failure
- Individual has sensorimotor or autonomic neuropathy not related to hATTR amyloidosis (including but not limited to, monoclonal gammopathy, autoimmune disease)
- Individual is using in combination with Onpattro, Tegsedi, Vyndaqel or Vyndamax
- Product use for non-FDA approved indications or indications not supported by industry-accepted guidelines.
- Doses, durations, or dosing intervals that exceed FDA maximum limits for any FDA-approved indication or are not supported by industry-accepted practice guidelines or peer-reviewed literature for the relevant off-label use.

Initial authorization is up to 12 months.

Deleted codes and codes which are not effective at the time the service is rendered may not be eligible for reimbursement.

U.S. Food and Drug Administration:

This section is to be used for informational purposes. FDA approval alone is not a basis for coverage.

Amvuttra[®] is a transthyretin-directed small interfering RNA indicated for the treatment of the polyneuropathy of hereditary transthyretin-mediated amyloidosis in adults.

References:

1. Lexi-Comp ONLINE™ with AHFS™, Hudson, Ohio: Lexi-Comp, Inc.; 2022; Updated periodically.
2. Amvuttra[®] [Prescribing Information]. Cambridge, MA: Alnylam Pharmaceuticals, Inc.

Date	Summary of Changes
March 2023	Criteria for use summary developed by the Ascension Medical Specialty Prior Authorization Team.
July 2023	Criteria for use summary approved by the Ascension Ambulatory Care Expert Review Panel.
August 2023	Criteria for use summary approved by the Ascension Therapeutic Affinity Group.

If you have questions, call [833-980-2352](tel:833-980-2352) to speak to a member of the Ascension Rx prior authorization team, or email your questions to smarthealthspecialty@ascension.org.