

## Amvuttra® (vutrisiran) Injectable **Medication Precertification Request**

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(All fields must be completed and legible for precertification review.)

**Aetna Precertification Notification** Phone: <u>1-866-752-7021</u> (TTY: 711)

1-888-267-3277

For Medicare Advantage Part B: Please Use Medicare Request Form

Please indicate:	☐ Start of treatment: Start of ☐ Continuation of therapy: I		_ ,	,					
Procertification	Requested By:				- nο·		Fax		
A. PATIENT INFO				1 110			I ax		
First Name:	MATION	Last Name:					DOB:		
Address:		Last Hame.	Ci	ty:			State:	ZIP:	
	WI-Di-			-				ZIF.	
Home Phone:	Work Pho			ell Phone:			Email:		
	Veight: lbs or kg	s Patient Height:	incl	nes or	_ cms   A	llergies:			
B. INSURANCE II									
Aetna Member ID		Does patient have other coverage?							
-			If yes, provide ID#: Carrier Name:						
Insured:		Insured:							_
Medicare: 🗌 Yes	s ☐ No If yes, provide ID #:		Med	icaid: 🗌 Yes	s □ No	If yes, p	rovide ID #:		
C. PRESCRIBER	INFORMATION								
First Name:		Last Name:				(Check O	<i>ne):</i> 🔲 M.D	. 🗌 D.O. 🗌 N.P. 🔲	P.A.
Address:				City:			State:	ZIP:	
Phone:	Fax:	St Lic #:		NPI #:		DEA #:		UPIN:	
Provider E-mail:	<u> </u>	Office Contact Na	ame:			•	Phor	ne:	
Specialty (Check	one): Neurologist Oth	er:					I		
	PROVIDER/ADMINISTRATION INF	·							
Place of Adminis				Dispensing	Provider	/Pharmac	v: (Patient s	elected choice)	
☐ Self-administe	ered Physician's Office	e		Dispensing Provider/Pharmacy: (Patient selected choice)  ☐ Physician's Office ☐ Retail Pharmacy					
		Specialty Pharmacy							
Center Name:					Name:				
Agency 1	n Center Phone: Name:								
☐ Administration		Diament Farm							
Address:				TIN:			PIN:		
E. PRODUCT INF	ORMATION								
Request is for: A	mvuttra (vutrisiran): Dose:			Frequency	:				
F. DIAGNOSIS IN	FORMATION – Please indicate prin	mary ICD Code and spec	ify any	other where ap	plicable.				
Primary ICD Cod	le: S	econdary ICD Code: _			c	ther ICD	Code:		
G. CLINICAL INFO	ORMATION – Required clinical info	rmation must be complet	ed in its	entirety for all	precertific	cation requ	ests.		
	(clinical documentation required)								
	s this infusion request in an outpatie								
└────────────────────────────────────									
(anaphylaxis, anaphylactoid reactions, myocardial infarction, thromboembolism, or seizures) during or immediately after									
administration?									
Yes No Does the patient have significant behavioral issues and/or physical or cognitive impairment that would impact the safety of the infusion therapy AND the patient does not have access to a caregiver?									
Please provide a description of the behavioral issue or impairment:									_
Yes No Is the patient medically unstable which may include respiratory, cardiovascular, or renal conditions that may limit the patient's ability to tolerate a large volume or load or predispose the patient to a severe adverse event that cannot be									
managed in an alternate setting without appropriate medical personnel and equipment?									
Please provide a description of the condition:   Cardiopulmonary:									
☐ Respiratory: ☐ Renal:									
				Other:					_
	Does the patient have a diagnosis o amyloid polyneuropathy [ATTR-FAF		ditary tr	ansthyretin-me	ediated am	yloidosis (	transthyretin-	type familial	
☐ Yes ☐ No \	·								
(	(e.g., amyloid deposition in biopsy specimens, TTR protein variants in serum, progressive peripheral sensory-motor polyneuropathy)?								
☐ Yes ☐ No I	s the patient a liver transplant recip	ient?							



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D (' ) F' (N			D # 1000							
Patient First Name	Patient Last Name	Patient Phone	Patient DOB							
G. CLINICAL INFORMATION – (continue	ed) Required clinical information must be completed	in its entirety for all precertification re	equests.							
Yes No Will the requested medication be used in combination any other medication approved for the treatment of hereditary transthyretin-mediated amyloidosis (e.g., Onpattro, Tegsedi, Vyndamax, Vyndaqel, Wainua)?										
☐ Yes ☐ No Will the requested medication be prescribed by or in consultation with any of the following: a) Neurologist, b) Geneticist, or c) Physician specializing in the treatment of amyloidosis?										
Yes DNo Has the patient experienced an ineffective response with Onpattro (patisiran)?										
Yes No Has the patient experienced an intolerance to Onpattro (patisiran)?										
Yes No Does the patient have a contraindication to Onpattro (patisiran)?										
For Continuation Requests (clinical documentation required):										
Yes No Has the patient demonstrated a beneficial response to the requested medication therapy compared to baseline (e.g., improvement of neuropathy severity and rate of disease progression as demonstrated by the modified Neuropathy Impairment Scale+7 (mNIS+7) composite score, the Norfolk Quality of Life-Diabetic Neuropathy (QoL-DN) total score, polyneuropathy disability (PND) score, FAP disease stage, manual grip strength)?										
H. ACKNOWLEDGEMENT										
Request Completed By (Signature Required): Date:/										
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or										

deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading,

The plan may request additional information or clarification, if needed, to evaluate requests.

commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.