

♦ aetna® Viltepso® (viltolarsen) Medication **Precertification Request**

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

FAX: <u>1-888-267-3277</u>

For Medicare Advantage Part B:

	(All fields must be	e completed and legible fo	r precertification rev	iew.)	Please Use M	edicare Request Form
Please indicate:	☐ Start of treatment: Start date	e <u> </u>				
	☐ Continuation of therapy: Da	te of last treatment	1 1			
Precertification R	equested By:		Phone	e:	Fax:	
A. PATIENT INFOR	RMATION					
First Name:		L	ast Name:			
Address:		C	City:		State:	ZIP:
Home Phone:	W	ork Phone:		Cell Phone:		
DOB:	Allergies:			Email:		
	lbs orkgs	Height:	inches	orcms		
B. INSURANCE INI		1101g11t				
	#:	Does patient have o	ther coverage?	☐ Yes ☐ No		
		If yes, provide ID#: _	_			
		Insured:				
Medicare: ☐ Yes	☐ No If yes, provide ID #:		Medicaid: ☐ Yes	☐ No If yes, pro	vide ID #:	
C. PRESCRIBER IN				<u> </u>		
First Name:		Last Name:		(Check One	e): 🔲 M.D. 🔲	D.O. 🗌 N.P. 🔲 P.A.
Address:		<u> </u>	City:		State:	ZIP:
Phone:	Fax:	St Lic #:	NPI #:	DEA #:	1	 JPIN:
Provider Email:		Office Contact Name			Phone:	·
	one): Neurologist Other:				1,	
	ROVIDER/ADMINISTRATION INFOR					
Place of Administ		WII/THOM	Dispensing F	Provider/Pharmacy	: Patient Sele	cted choice
☐ Self-administer				-	Retail Pharm	
Outpatient Infus			-		_	
	ime:		- Name:			
Home Infusion			· · · · · · · · · · · · · · · · · · ·			
Address:	ame:					
	code(s) (CPT):					
E. PRODUCT INFO						
	tepso (viltolarsen) Dose:		F	requency:		
-	ORMATION – Please indicate primar	ry ICD Code and specify a				
Primary ICD Code		condary ICD Code:	, , , , , , , , , , , , , , , , , , , ,		ode:	
<u> </u>	RMATION – Required clinical inform				·	
	clinical documentation required):	•		•		
☐ Yes ☐ No Is t	his infusion request in an outpatient h					
	Yes No Has the patient experi					
	\ 0 /	etaminophen, steroids, dip (anaphylaxis, anaphylacto	,	· ·		,
	immediately after an ir	nfusion?		·	•	,
	Yes No Does the patient have outpatient hospital set		sues that require the	use of special interve	entions only ava	ailable in the
Ę	Yes No Does the patient have	significant behavioral issu			nt that would im	pact the safety of the
		the patient does not have scription of the behavioral i				
	Yes No Is the patient medically				onditions that n	nay limit the
T	patient's ability to toler	rate a large volume or load	d or predispose the p	patient to a severe ad	verse event tha	t cannot be
		ate setting without appropr scription of the condition: [
	,					
			Renal:			
			Other:			
	es the patient have a documented dia	agnosis of Duchenne musc	cular dystrophy (DM	ע)?		

☐ Yes ☐ No Will the medication be prescribed by or in consultation with a physician who specializes in the treatment of Duchenne muscular dystrophy (DMD)?



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Page 2 of 2

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

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continued) - F	Required clinical information must be compl	eted in its entirety for all precertifi	cation requests					
Yes No Will the requested medication Yes No Does the patient's dose excee	be used concomitantly with golodirsen? d 80 mg/kg once weekly?	eted in its <u>criticity</u> for an precentin	Sauon requests.					
For Initiation of Therapy (clinical documentation required):								
☐ Yes ☐ No Was genetic testing conducted to confirm the diagnosis of Duchenne muscular dystrophy (DMD)? ☐ Yes ☐ No Was genetic testing conducted to identify the specific type of DMD gene mutation?								
Please indicate the DMD gene mutation:								
☐ Yes ☐ No Is the DMD gene mutation amenable to exon 53 skipping?								
☐ Yes ☐ No Is the patient able to walk independently without assistive devices?								
☐ Yes ☐ No Will treatment with the requested medication be initiated prior to age 10?								
Yes No Has the patient previously received gene replacement therapy for DMD (e.g., Elevidys)?								
	nt experienced a worsening in clinical status MD (e.g., Elevidys)?	s (e.g., decline in ambulatory fund	tion) since receiving gene replacement					
For patients re-starting therapy with the requested medication after administration of gene therapy (clinical documentation required):								
☐ Yes ☐ No Was genetic testing conducted ☐ Yes ☐ No Was genetic testing conducted ☐ Please indicate the DMD general conducted the DMD general condu	I to identify the specific type of DMD gene i	, , ,						
☐ Yes ☐ No Is the DMD gene mutation amenable to exon 53 skipping?								
☐ Yes ☐ No Is the patient able to walk independently without assistive devices?								
☐ Yes ☐ No Will treatment with the requested medication be initiated prior to age 10?								
Yes No Has the patient experienced a DMD (e.g., Elevidys)?	worsening in clinical status (e.g., decline in	ambulatory function) since recei	ving gene replacement therapy for					
For Continuation of Therapy (clinical documentation required):								
☐ Yes ☐ No Has the patient demonstrated	a response to therapy as evidenced by rem	naining ambulatory (e.g., not whee	elchair dependent)?					
H. ACKNOWLEDGEMENT								
Request Completed By (Signature Require	red):		Date: / /					
Any person who knowingly files a request fo insurance company by providing materially insurance act, which is a crime and subjects	false information or conceals materia	information for the purpose of						

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