SENTARA COMMUNITY PLAN (MEDICAID)

MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions:</u> The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; <u>fax to 1-844-305-2331</u>. No additional phone calls will be necessary if all information (<u>including phone and fax #s</u>) on this form is correct. <u>If information provided is not complete, correct, or legible, authorization can be delayed</u>.

<u>Drug Requested</u>: Elevidys (delandistrogene moxeparvovec-rokl) J1413 (MEDICAL)

Member Name:	
Member Sentara #:	
Prescriber Name:	
	Date:
Office Contact Name:	
Pnone Number:	Fax Number:
DEA OR NPI #: DRUG INFORMATION: Author	
DEA OR NPI #: DRUG INFORMATION: Author Drug Form/Strength:	orization may be delayed if incomplete.
DEA OR NPI #: DRUG INFORMATION: Author Drug Form/Strength: Dosing Schedule:	orization may be delayed if incomplete.
DEA OR NPI #: DRUG INFORMATION: Author Drug Form/Strength: Dosing Schedule:	Derization may be delayed if incomplete. Length of Therapy: ICD Code, if applicable:

Recommended Dose: The recommended dose of Elevidys is 1.33×10^{14} vector genomes per kilogram (vg/kg) of body weight (or 10 mL/kg body weight). For the number of vials required, refer to table below. Calculate the dose as follows: ELEVIDYS dose (in mL) = patient body weight (in kilogram) x 10 The multiplication factor 10 represents the per kilogram dose $(1.33 \times 10^{14} \text{ vg/kg})$ divided by the amount of vector genome copies per mL of the ELEVIDYS suspension $(1.33 \times 10^{13} \text{ vg/mL})$. Number of ELEVIDYS vials needed = ELEVIDYS dose (in mL) divided by 10 (round to the nearest number of vials).

Patient Weight (kg)	Total Vials per Kit	Total Dose Volume per Kit (mL)	NDC Number
10.0 - 10.4	10	100	60923-501-10
10.5 – 11.4	11	110	60923-502-11
11.5 – 12.4	12	120	60923-503-12
12.5 – 13.4	13	130	60923-504-13
13.5 – 14.4	14	140	60923-505-14
14.5 – 15.4	15	150	60923-506-15
15.5 – 16.4	16	160	60923-507-16
16.5 – 17.4	17	170	60923-508-17
17.5 – 18.4	18	180	60923-509-18
18.5 – 19.4	19	190	60923-510-19
19.5 – 20.4	20	200	60923-511-20
20.5 – 21.4	21	210	60923-512-21
21.5 – 22.4	22	220	60923-513-22
22.5 – 23.4	23	230	60923-514-23
23.5 – 24.4	24	240	60923-515-24
24.5 – 25.4	25	250	60923-516-25
25.5 – 26.4	26	260	60923-517-26
26.5 – 27.4	27	270	60923-518-27
27.5 – 28.4	28	280	60923-519-28
28.5 – 29.4	29	290	60923-520-29
29.5 – 30.4	30	300	60923-521-30
30.5 – 31.4	31	310	60923-522-31
31.5 – 32.4	32	320	60923-523-32
32.5 – 33.4	33	330	60923-524-33
33.5 – 34.4	34	340	60923-525-34
34.5 – 35.4	35	350	60923-526-35
35.5 – 36.4	36	360	60923-527-36
37.5 – 38.4	38	380	60923-529-38
38.5 – 39.4	39	390	60923-530-39
39.5 – 40.4	40	400	60923-531-40

Patient Weight (kg)	Total Vials per Kit	Total Dose Volume per Kit (mL)	NDC Number
40.5 – 41.4	41	410	60923-532-41
41.5 – 42.4	42	420	60923-533-42
42.5 – 43.4	43	430	60923-534-43
43.5 – 44.4	44	440	60923-535-44
44.5 – 45.4	45	450	60923-536-45
45.5 – 46.4	46	460	60923-537-46
46.5 – 47.4	47	470	60923-538-47
47.5 – 48.4	48	480	60923-539-48
48.5 – 49.4	49	490	60923-540-49
49.5 – 50.4	50	500	60923-541-50
50.5 – 51.4	51	510	60923-542-51
51.5 – 52.4	52	520	60923-543-52
52.5 – 53.4	53	530	60923-544-53
53.5 – 54.4	54	540	60923-545-54
54.5 – 55.4	55	550	60923-546-55
55.5 – 56.4	56	560	60923-547-56
56.5 – 57.4	57	570	60923-548-57
57.5 – 58.4	58	580	60923-549-58
58.5 – 59.4	59	590	60923-550-59
59.5 – 60.4	60	600	60923-551-60
60.5 – 61.4	61	610	60923-552-61
61.5 – 62.4	62	620	60923-553-62
62.5 – 63.4	63	630	60923-554-63
63.5 – 64.4	64	640	60923-555-64
64.5 – 65.4	65	650	60923-556-65
65.5 – 66.4	66	660	60923-557-66
66.5 – 67.4	67	670	60923-558-67
67.5 – 68.4	68	680	60923-559-68
68.5 – 69.4	69	690	60923-560-69
69.5 and above	70	700	60923-561-70

CLINICAL CRITERIA: Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

Authorization: One treatment per lifetime or until 6 years of age, whichever is first

Member is between 4 and 5 years of age
Member is <u>NOT</u> on concomitant therapy with DMD-directed antisense oligonucleotides and will stop therapy prior to Elevidys administration (e.g., golodirsen, casimersen, viltolarsen, eteplirsen)
Member has a NOT received a DMD-directed antisense oligonucleotide within the past 30 days
Member does NOT have an active infection, including clinically important localized infections
Member has been on a stable dose of a corticosteroid, unless contraindicated or intolerance, prior to start of therapy and will be used concomitantly with a corticosteroid regimen pre- and post- infusion (refer to the package insert for recommended corticosteroid dosing during therapy; verified by chart notes and/or pharmacy paid claims)
Member's troponin-I levels will be monitored at baseline and subsequently as clinically indicated
Provider will submit member's baseline liver function assessed 30 days prior to request and following therapy for at least 3 months and as indicated (submit documentation; Provider please note: Requested medication will not be approved if GGT is < 3 times upper limit of normal and Total bilirubin is < 3 times the upper limit of normal)
Member has a confirmed mutation of the DMD gene between exons 18-58 (submit genetic testing)
Member is ambulatory as confirmed by the North Star Ambulatory Assessment (NSAA) scale (i.e., patient score of 1 or greater for each question) (submit scale)
Member is receiving physical and/or occupational therapy
Member must have a baseline anti-AAVrh74 total binding antibody titer of < 1:400 as measured by ELISA (submit analysis)
Member does NOT have any deletion in exon 8 and/or exon 9 in the DMD gene
Member has <u>NOT</u> had gene therapy, cell-based therapy, or clustered regularly interspaced short palindromic repeats (CRISPR/Cas9)

Medication being provided by: Please check applicable box below.				
	Location/site of drug administration:			
	NPI or DEA # of administering location:			
	<u>OR</u>			
	Specialty Pharmacy – Proprium Rx			

For urgent reviews: Practitioner should call Sentara Health Pre-Authorization Department if they believe a standard review would subject the member to adverse health consequences. Sentara Health's definition of urgent is a lack of treatment that could seriously jeopardize the life or health of the member or the member's ability to regain maximum function.

**Use of samples to initiate therapy does not meet step edit/ preauthorization criteria. ** *Previous therapies will be verified through pharmacy paid claims or submitted chart notes. *

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REVISED/UPDATED: 9/11/2023;10/17/2023

^{*}Approved by Pharmacy and Therapeutics Committee: 9/21/2023