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 (including phone and fax #s) on this form is correct. <u>If information provided is not complete, correct, or legible, authorization can be delayed</u>.

Drug Requested: Aldurazyme® (laronidase) IV solution (J1931) (Medical)

MEMBER & PRESCRIBER INFORM	ATION: Authorization may be delayed if incomplete.	
Member Name:		
Member Sentara #:	Date of Birth:	
Prescriber Name:		
Prescriber Signature:		
Office Contact Name:		
Phone Number: Fax Number:		
DEA OR NPI #:		
DRUG INFORMATION: Authorization m	ay be delayed if incomplete.	
Drug Form/Strength:		
Dosing Schedule:	Length of Therapy:	
Diagnosis:	ICD Code, if applicable:	
Weight (current):	Weight (within last 30 days):	
Quantity Limit (max daily dose) [NDC unit]: 2.9mg vial; 92 vials every 28 days		
Max Units (per dose and over time) [HCPCS unit]: 667 billable units every 7 days		
	meframe does not jeopardize the life or health of the member unction and would not subject the member to severe pain.	
	hat apply. All criteria must be met for approval. To luding lab results, diagnostics, and/or chart notes, must	
Initial Approval Authorization – 6 mont	ths	

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 \square Member is ≥ 6 months of age

	M	such as has a definitive diagnosis of MDS I confirmed by one of the fellowing.
		ember has a definitive diagnosis of MPS I confirmed by one of the following:
		Detection of biallelic pathogenic mutations in the IDUA gene by molecular genetic testing
		Fibroblast or leukocyte alpha-L-iduronidase (IDUA) enzyme activity level of less than 10% of the lower limit of the normal range of the measuring laboratory
	Me	ember has diagnosis of one of the following:
		Diagnosis of Hurler (severe) or Hurler-Scheie (attenuated) forms of disease
		Diagnosis of Scheie (attenuated) form of disease with moderate to severe symptoms
	Me	ember has absence of severe cognitive impairment
	Documented baseline value for urinary glycosaminoglycan (uGAG)	
	pre wa mi	cumented baseline values for one or more of the following \square Members 6 years or greater: percent edicted forced vital capacity (FVC) of \leq 77% of the patient's predicted normal FVC value, 6-minute lk test (must be able to stand independently for 6 minutes and walk a minimum of 5 meters within 6 nutes), joint range of motion, left ventricular hypertrophy, growth, quality of life (CHAQ/HAQ/MPS AQ);
		OR
		Members 6 months to less than 6 years: cardiac status, upper airway obstruction during sleep, growth velocity, mental development, FVC, and/or 6-minute walk test (must be able to stand independently for 6 minutes and walk a minimum of 5 meters within 6 minutes)
nt	inu	ation of Therapy – 12 month Approval
	Me	ember continues to meet all initial authorization criteria
	Member has absence of unacceptable toxicity from the drug. Examples of unacceptable toxicity include: anaphylaxis and severe hypersensitivity reactions, acute respiratory complications, acute cardiorespiratory failure, severe infusion reactions, etc	
	Me	ember does not have progressive/irreversible severe cognitive impairment.
	Me	ember has a documented reduction in uGAG levels compared to pretreatment baseline
		ember has demonstrated a beneficial response to therapy compared to pretreatment baseline in one or ore of the following:
		Members 6 years or greater: stability or improvement in percent predicted FVC and/or 6- minute walk test, increased joint range of motion, decreased left ventricular hypertrophy, improved growth, improved quality of life (clinically meaningful change in the CHAQ/HAQ/MPS HAQ disability index);
		OR
		Members 6 months to less than 6 years: stability or improvement in cardiac status, upper airway obstruction during sleep, growth velocity, mental development, FVC and/or 6-minute walk test

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