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>: ELAPRASE[®] (idursulfase) (IV INFUSION ONLY) (J1743) (Medical)

MEMBER & PRESCRIBER INFO	RMATION: Authorization may be delayed if incomplete.
Member Name:	
Member Sentara #:	Date of Birth:
Prescriber Name:	
Prescriber Signature:	Date:
Office Contact Name:	
Phone Number:	Fax Number:
DEA OR NPI #:	
DRUG INFORMATION: Authorizati	on may be delayed if incomplete.
Drug Form/Strength:	
Dosing Schedule:	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight (current):	Weight (within last 30 days):
	the timeframe does not jeopardize the life or health of the member am function and would not subject the member to severe pain.
	all that apply. All criteria must be met for approval. To a, including lab results, diagnostics, and/or chart notes, must
Initial Approval Authorization – 6 n	nonths. (Max approved dose will be 0.5mg/kg every 7 days)
\square Member is ≥ 5 years of age	
☐ Provider is a specialist in genetics or i	metabolic disorders
☐ Member has absence of severe cognit	ive impairment
 Patient has a diagnosis of Hunter dise 	ase (also referred to as Mucopolysaccharidosis II; MPS II)

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_	Diag	gnosis of Hunter disease has been confirmed by one of the following:
		Deficient iduronate 2-sulfatase (I2S) enzyme activity in white cells, fibroblasts, or plasma in the presence of normal activity of at least one other sulfatase;
		OR
		Detection of pathogenic mutations in the IDS gene by molecular genetic testing
	Doc	umented baseline value for urinary glycosaminoglycan (uGAG)
	Doc	umented baseline values for at least one of the following:
		Member ≥ 5 years of age: 6-minute walk test (6-MWT) and/or percent predicted forced vital capacity (FVC)
		OR
		Member < 5 years of age: spleen volume; liver volume; FVC; and/or 6-minute walk test
		SION CRITERIA: Elaprase® is considered investigational when used for any
indic	atio	n not listed above.
Cont	inua	tion of Therapy – 6 months Approval (Max dose 60 billable units every 7 days)
	Prov	vider is a specialist in genetics or metabolic disorders, a cardiologist, or a nephrologist
	Mer	mber continues to meet the criteria in initial section
	hyp	ence of unacceptable toxicity from the drug. Examples include the following: severe ersensitivity including anaphylactic and anaphylactoid reactions; antibody development and serious erse reactions; acute respiratory complications; acute cardiorespiratory failure; etc.
		sise reactions, acute respiratory complications, acute caratorespiratory range, etc.
	Mer	nber does not have progressive/irreversible severe cognitive impairment
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_	Mer Mer	mber does not have progressive/irreversible severe cognitive impairment
	Mer Mer mor	mber does not have progressive/irreversible severe cognitive impairment mber has documented reduction in uGAG levels mber has demonstrated beneficial response to therapy compared to pretreatment baseline in one or
	Mer Mer mor	mber does not have progressive/irreversible severe cognitive impairment mber has documented reduction in uGAG levels mber has demonstrated beneficial response to therapy compared to pretreatment baseline in one or e of the following
	Mer mor	mber does not have progressive/irreversible severe cognitive impairment mber has documented reduction in uGAG levels mber has demonstrated beneficial response to therapy compared to pretreatment baseline in one or e of the following Members ≥5 years: stabilization or improvement in 6-MT and/or FVC

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limited studies on members with severe cognitive impairment.

Medic	cation being provided by (check applicable box below):
	Location/site of drug administration:
]	NPI or DEA # of administering location:
	OR
	Specialty Pharmacy - PropriumRx
review v treatmen	ent reviews: Practitioner should call Sentara Pre-Authorization Department if they believe a standard would subject the member to adverse health consequences. Sentara's definition of urgent is a lack of nt that could seriously jeopardize the life or health of the member or the member's ability to regain um function.
** U	se of samples to initiate therapy does not meet step edit/preauthorization criteria.**
* <u>Previ</u>	ious therapies will be verified through pharmacy paid claims or submitted chart notes.*