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>: Nulibry[™] (fosdenopterin) IV (Medical) (J3490)

MEMBER & PRESCRIBER INFORMATION: 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: Author	rization may be delayed if incomplete.
Drug Form/Strength:	
Dosing Schedule:	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight:	Date:
	ox, the timeframe does not jeopardize the life or health of the member timum function and would not subject the member to severe pain.
Quantity Limit: Maximum approval	of 0.9mg/kg/day (actual body weight)
	for infants will be 0.55mg/kg/dose once daily for 1 month, then r 2 months, then increase to target dose of 0.9mg/kg once daily
	elow all that apply. All criteria must be met for approval. To ation, including lab results, diagnostics, and/or chart notes, must be
Initial Approval : 6 months	

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Provider is a metabolic geneticist, neurologist, or other specialist in treatment of molybdenum cofactor

deficiency (MoCD) Type A

u	approved test documenting a mutation in the MOCS1 gene (must submit genetic test results)
	Member has confirmation of all of the following (must submit lab test results):
	☐ Elevated S-sulfocysteine or sulfite urinary levels
	☐ Low serum or urinary uric acid levels
	☐ Elevated xanthine or hypoxanthine urinary levels
	Member has clinical presentation of MoCD including at least two (2) of the following (submit current chart documentation):
	□ intractable seizures
	encephalopathy
	□ hyper/hypotonia, feeding difficulties
	□ developmental delay
	□ exaggerated startle reaction
	Member's current weight must be noted: (submit current chart notes documenting weight)
	Was member already initiated on fosdenopterin (Nulibry) or on recombinant cPMP (rcPMP)?
	☐ Yes (must submit chart note documentation)
	□ No
	Member will not use fosdenopterin in combination with other substrate replacement therapy (e.g., recombinant cyclic pyranopterin monophosphate, etc.)
	Member does not have clinically significant intracranial hemorrhage, cortical or subcortical encephalomalacia, or abnormalities on brain imaging not attributable to MoCD Type A
	Member does not have a Modified Glasgow Coma Scale (mGCS) for infants and children score of less than 7 for more than 24 hours (must submit mGCS scale with results)
appro	thorization Approval – 12 months: Check below all that apply. All criteria must be met for val. To support each line checked, all documentation, including lab results, diagnostics, and/or chart must be provided or request may be denied.
	If established on Nulibry but not previously approved by Sentara Health Plans <u>ALL</u> of the initial authorization criteria must be met
	Member has confirmation of both of the following (must submit lab test results):
	☐ Reduction of S-sulfocysteine (SSC) urinary levels to ≤11 μmol/mmol
	☐ Serum or urinary uric acid levels have increased from baseline or have been maintained above baseline level since last approval
	Member has had stabilization or improvement in one or more signs and symptoms of disease including, but not limited to, seizure frequency/duration, growth, achievement of developmental milestones

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PA Nulibry (Medical) (Medicaid) (continued from previous page)

	Member's current weight must be noted: (submit current chart notes documenting weight)
	Member does not have a Modified Glasgow Coma Scale (mGCS) for infants and children score of less than 7 for more than 24 hours (must submit mGCS scale with results)
Medic	ation being provided by: Please check applicable box below.
□ Lo	ocation/site of drug administration:
N	PI or DEA # of administering location:
	<u>OR</u>
□ Sp	ecialty Pharmacy – PropriumRx
standard is a lack	ent reviews: Practitioner should call Sentara Health Pre-Authorization Department if they believe a dreview would subject the member to adverse health consequences. Sentara Health's definition of urgent to of treatment that could seriously jeopardize the life or health of the member or the member's ability to naximum function.
**	Use of samples to initiate therapy does not meet step edit/preauthorization criteria.**
Prev	ious therapies will be verified through pharmacy paid claims or submitted chart notes.