## 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: Evkeeza® (evinacumab-dgnb) (J1305) (Medical)

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.					
Member Name:					
Member Sentara #:	Date of Birth:				
Prescriber Name:					
	Date:				
Phone Number:					
DEA OR NPI #:					
DRUG INFORMATION: Author	orization may be delayed if incomplete.				
Drug Form/Strength:					
Dosing Schedule:	Length of Therapy:				
Diagnosis:	ICD Code, if applicable:				
Weight:	Date:				
☐ Standard Review. In checking this	box, the timeframe does not jeopardize the life or health of the member aximum function and would not subject the member to severe pain.				

## Quantity Limit (max daily dose) [NDC Unit]:

- Evkeeza 345 mg/2.3 mL single-dose vial: 2 vials per 28 days
- Evkeeza 1200 mg/8 mL single-dose vial: 1 vial per 28 days

## Max Units (per dose and over time) [HCPCS Unit]:

• 1690 mg every 28 days

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**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.

**Initial Authorization: 6 months** 

Member is 5 years of age or older				
Baseline low-density lipoprotein cholesterol (LDL-C), total cholesterol (TC), apolipoprotein B (apo B) and non-high density lipoprotein cholesterol (non-HDL-C) labs must be obtained prior to initiating treatment (please submit labs with request)				
Member does <b>NOT</b> have heterozygous familial hypercholesterolemia (HeFH)				
Requested medication is prescribed by or in consultation with a specialist in cardiology, lipidology, or endocrinology				
Member has a diagnosis of Homozygous Familial Hypercholesterolemia (HoFH) confirmed by at least <b>ONE</b> of the following:				
□ Documented DNA test for functional mutation(s) in LDL receptor alleles or alleles known to affect LDL receptor functionality (submit test results with request)				
□ Untreated LDL-C > 500 mg/dL <b>OR</b> treated LDL-C $\geq$ 300 mg/Dl along with <b>ONE</b> of the following				
☐ Cutaneous or tendon xanthoma before 10 years of age				
<ul> <li>Untreated LDL cholesterol levels consistent with heterozygous FH in both parents (&gt; 190 mg/dL)</li> </ul>				
Member has been receiving stable background lipid lowering therapy for at least 4 weeks (verified by pharmacy paid claims)				
Requested therapy will be used in conjunction with a low-fat or heart-healthy diet and other LDL-lowering therapies (e.g., statins, ezetimibe, PCSK9 inhibitors, Juxtapid, LDL apheresis)				
Member has had an unsuccessful 3-month trial of <u>ALL</u> the following (verified by pharmacy paid claims):				
<ul> <li>□ Highest available (or maximally tolerated) dose of atorvastatin OR rosuvastatin</li> <li>□ ezetimibe (Zetia<sup>®</sup>)</li> </ul>				
□ PCSK9 inhibitor indicated for HoFH (e.g., Repatha), unless contraindicated				
Despite pharmacological treatment with a PCSK9 inhibitor, maximally tolerated statin therapy, and ezetimibe, the member's LDL cholesterol $\geq 100$ mg/dL (or $\geq 70$ mg/dL for members with clinical atherosclerotic cardiovascular disease [ASCVD]) while on therapy				

\*Provider please note: If the member is not able to use the maximum dose of atorvastatin or rosuvastatin due to muscle symptoms, documentation of a causal relationship must be established between statin use and muscle symptoms

	Me	ember m	nust meet <b>ONE</b> of the following:		
			al documentation must demonstrate that the member experienced pain, tenderness, stiffness,		
		-	ng, weakness, and/or fatigue in addition to <u>ALL</u> the following:		
			sscle symptoms resolve after discontinuation of statin		
		□ Mu	scle symptoms occurred when re-challenged at a lower dose of the same statin		
		□ Mu	sscle symptoms occurred after switching to an alternative statin		
		ren	cumentation ruling out non-statin causes of muscle symptoms (e.g., hypothyroidism, reduced al function, reduced hepatic function, rheumatologic disorders, such as polymyalgia rumatica, steroid myopathy, vitamin D deficiency, or primary muscle disease)		
		suppor	er has been diagnosed with rhabdomyolysis associated with statin use and the diagnosis is ted by acute neuromuscular illness or dark urine $\underline{AND}$ an acute elevation in creatine kinase $y > 5,000 \text{ IU/L}$ or 5 times the upper limit of normal [ULN])		
Rea	uth	orizati	on: 12 months. 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.					
		-	therapy continues to be used in conjunction with a low-fat or heart-healthy diet and other ring therapies (e.g., statins, ezetimibe, PCSK9 inhibitors, Juxtapid, LDL apheresis)		
☐ Member has experienced an absence of unacceptable toxicity from therapy (e.g., severe hypersensitivity)					
☐ Member has had a reduction in LDL-C when compared to the initial baseline labs (please submit labs collected within the last 30 days)					
Me	dica	ation b	eing provided by: Please check applicable box below.		
	Loca	ation/sit	te of drug administration:		
	NPI or DEA # of administering location:				
OR					
	□ Specialty Pharmacy – PropriumRx				

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. \*