SENTARA HEALTH PLANS

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid)</u> on this request. All other information may be filled in by office staff; <u>fax to 1-800-750-9692</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 the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

<u>Drug Requested</u> : (select drug below)					
□ Repatha [®] (evolocumab)	□ Praluent® (alirocumab)				
MEMBER & PRESCRIBER INFORMA	ATION: Authorization may be delayed if incomplete.				
Member Name:					
Member Sentara #:					
Prescriber Name:					
Prescriber Signature:					
Office Contact Name:					
Phone Number:					
DEA OR NPI #:					
DRUG INFORMATION: Authorization ma	ay be delayed if incomplete.				
Drug Form/Strength:					
Dosing Schedule:	Length of Therapy:				
Diagnosis:					
Weight:					
DRUG	QUANTITY LIMIT				
PRALUENT 150 MG/ML PEN	2 pens per 28 days				
PRALUENT 75 MG/ML PEN	2 pens per 28 days				
REPATHA 140 MG/ML SURECLICK	2 auto-injectors per 28 days				
REPATHA 140 MG/ML SYRINGE	2 syringes per 28 days				
REPATHA 420 MG/3.5ML PUSHTRONX	1 cartridge per 28 days				
	hat apply. All criteria must be met for approval. To luding lab results, diagnostics, and/or chart notes, must be				

Initial Authorization: 12 months

	Must be prescribed by or in consultation with a Cardiologist, Endocrinologist or Lipid Specialist
	Medication will be used as adjunct to low-fat diet
	Provider has COMPLETED Sections I, IIa or IIb and III (if applicable) below
Sect	tion I. Diagnoses: (select one below)
	Diagnosis: Primary Hyperlipidemia
hyper as co	<u>CE</u> : This is not associated with atherosclerotic cardiovascular disease (ASCVD), heterozygous familial rcholesterolemia (HeFH), or homozygous familial hypercholesterolemia (HoFH) and may be referred to mbined hyperlipidemia, hypercholesterolemia (pure, primary), dyslipidemia, or increased/elevated low-ty lipoprotein cholesterol (LDL-C) levels.
	Member must meet <u>ALL</u> the following:
	☐ Member is 18 years of age or older
	\square Member has a coronary artery calcium or calcification score ≥ 300 Agatston units
	☐ Member meets <u>ONE</u> of the following:
	☐ Member meets <u>ALL</u> the following:
	 Member has tried one high-intensity statin therapy (i.e., atorvastatin ≥ 40 mg daily; rosuvastatin ≥ 20 mg daily [as a single-entity or as a combination product])
	\square Member has tried one high-intensity statin therapy above along with ezetimibe (as a single-entity or as a combination product) for ≥ 8 continuous weeks
	☐ Member's LDL-C level after this treatment regimen remains ≥ 100 mg/dL
	☐ Member has been determined to be statin intolerant and meets all clinical criteria in section IIb below
	Provider has completed section III if applicable
o D	Diagnosis: Atherosclerotic Cardiovascular Disease
	Member is 18 years of age or older and has Atherosclerotic Cardiovascular Disease (ASCVD) confirmed by at least ONE of the following:
	☐ Acute Coronary Syndrome
	☐ History of myocardial infarction
	□ Stable or unstable angina
	☐ Peripheral arterial disease presumed to be of atherosclerotic origin
	☐ Member has undergone coronary or other arterial revascularization procedure in the past
	☐ History of Stroke
	☐ History of Transient ischemic attack
	Provider has completed sections IIa or IIb & III if applicable

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□ D	iag	nos	sis:	Heterozygous familial hypercholesterolemia (HeFH)
				s 10 years of age or older and has heterozygous familial hypercholesterolemia (HeFH) as by the following:
		Me	mb	er meets ONE of the following:
			Me	ember has an untreated low-density lipoprotein cholesterol (LDL-C) \geq 190 mg/dL (prior to atment with antihyperlipidemic therapy)
			the	ember has genetic confirmation of heterozygous familial hypercholesterolemia by mutations in low-density lipoprotein receptor, apolipoprotein B, proprotein convertase subtilisin kexin be 9, or low-density lipoprotein receptor adaptor protein 1 gene
				ember has been diagnosed with heterozygous familial hypercholesterolemia by meeting ONE the following diagnostic criteria thresholds:
				Provider attests member's Dutch Lipid Network criteria score was > 5
				Provider attests that Simone Broome criteria met the threshold for "definite" or "possible (or probable)" familial hypercholesterolemia
	Pro	ovid	er h	as completed sections IIa or IIb & III if applicable
□ D	iag	nos	sis:	Homozygous familial hypercholesterolemia (HoFH)
				s 10 years of age or older and has homozygous familial hypercholesterolemia (HoFH) as by the following:
		Me	mb	er meets ONE of the following:
			apo	ember has genetic confirmation of two mutant alleles at the low-density lipoprotein receptor olipoprotein B, proprotein convertase subtilisin kexin type 9 (PCSK9), or low-density oprotein receptor adaptor protein 1 gene locus
				ember has an untreated low-density lipoprotein cholesterol (LDL-C) level > 500 mg/dL AND tets \underline{ONE} of the following:
				Member has had clinical manifestations of homozygous familial hypercholesterolemia before the age of 10 (e.g., xanthomas, tendon xanthomas, arcus cornea, tuberous xanthomas, or xanthelasma)
				Members parents both have had untreated LDL-C levels or total cholesterol levels consistent with heterozygous familial hypercholesterolemia (i.e., both parents have had an untreated LDL-C level \geq 190 mg/dL and/or an untreated total cholesterol level $>$ 250 mg/dL
			Me	ember has a treated LDL-C level $\geq 300 \text{ mg/dL AND}$ meets <u>ONE</u> of the following:
				Member has had clinical manifestations of homozygous familial hypercholesterolemia before the age of 10 (e.g., xanthomas, tendon xanthomas, arcus cornea, tuberous xanthomas, or xanthelasma)
				Members parents both have had untreated LDL-C levels or total cholesterol levels consistent with heterozygous familial hypercholesterolemia (i.e., both parents have had an untreated LDL-C level $\geq 190~\text{mg/dL}$ and/or an untreated total cholesterol level $\geq 250~\text{mg/dL}$
	Pro	vid	er h	as completed sections IIa or IIb & III if applicable

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Sect thera		IIa. FOR ALL DIAGNO	OSIS: Skip to Section	IIb IF member is unable to tolerate statin
	Me		lowing statin therapies	as a single-entity or combination product for at
	lea	st 8 consecutive weeks:		
		High intensity statin therapy v	` -	
		High intensity statin therapy r	osuvastatin (generic C	restor) \geq 20 mg daily
		Moderate-intensity statin thera	apy (member unable to	tolerate high intensity therapy)
		Low intensity statin therapy (1	member unable to toler	rate moderate intensity therapy
	Me	ember's LDL-C after 8-week tr	rial of maximally tolera	ated statin therapy remains $\geq 70 \text{ mg/dL}$
	Ple	ease provide member's LDL lev	vels below:	
	LI	OL baseline:	LD	DL post-treatment:
Sect	tion	IIb. FOR ALL DIAGNO	OSIS: Contraindicatio	on to statin therapy
Ω.	elect	below if the member is unable	to tolerate low moder	rate, and high intensity statin therapy as
ev	iden		nt symptoms to TWO	different statins (i.e., more than 2 weeks); Please
ev pr	iden ovid	ced by intolerable and persister e previously attempted statin na	nt symptoms to <u>TWO</u> ame, strength & therap	different statins (i.e., more than 2 weeks); Please
ev pr D i	iden ovid rug	ced by intolerable and persister the previously attempted statin na Name:	nt symptoms to TWO ame, strength & therap <a "two<="" a="" href="Strength: " strength:="">	different statins (i.e., more than 2 weeks); Please by initiation date below:
ev pr D i	riden ovid rug rug Me	ced by intolerable and persister to previously attempted statin not name: Name: Name:	nt symptoms to TWO ame, strength & therap Strength: Strength:	different statins (i.e., more than 2 weeks); Please by initiation date below: Date started:
ev pr Di Di	riden ovid rug rug Me	ced by intolerable and persister te previously attempted statin no Name: Name: ember is unable to tolerate static	nt symptoms to TWO ame, strength & therap Strength: Strength: n therapy due to the oc	different statins (i.e., more than 2 weeks); Please by initiation date below: Date started: Date started:
ev pr Di Di	riden ovid rug rug Me	Name: ember is unable to tolerate static mptoms:	nt symptoms to TWO ame, strength & therap Strength: Strength: n therapy due to the occitiont CK elevations)	different statins (i.e., more than 2 weeks); Please by initiation date below: Date started: Date started: ccurrence of at least ONE of the following
ev pr Di Di	riden ovid rug rug Me syr	Name: ember is unable to tolerate static mptoms: Myalgia (muscle symptoms w Myositis (muscle symptoms w	nt symptoms to TWO ame, strength & therap Strength: Strength: n therapy due to the occ without CK elevations) with CK elevations < 10	different statins (i.e., more than 2 weeks); Please by initiation date below: Date started: Date started: ccurrence of at least ONE of the following
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ev pr Di Di	riden ovid rug rug Me syn	Name: ember is unable to tolerate statismptoms: Myalgia (muscle symptoms w Myositis (muscle symptoms w Member has experienced rhablimit of normal	nt symptoms to TWO ame, strength & therap Strength: Strength: n therapy due to the occ without CK elevations) with CK elevations < 10 odomyolysis or muscle	different statins (i.e., more than 2 weeks); Please by initiation date below:
ev pr Dr	riden ovid rug rug Me syn	Name: ember is unable to tolerate statismptoms: Myalgia (muscle symptoms w Myositis (muscle symptoms w Member has experienced rhablimit of normal Member has a labeled contrain	nt symptoms to TWO ame, strength & therap Strength: Strength: n therapy due to the occ without CK elevations) with CK elevations < 10 odomyolysis or muscle ndication to ALL static been attempted and fa	different statins (i.e., more than 2 weeks); Please by initiation date below:
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Reauthorization: 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.

<u>Provider please note</u>: a one-time reauthorization is required after initial 12-month approval

Provider attests member has experienced a positive clinical response to PCSK9 therapy (e.g., decreasing low-density lipoprotein cholesterol (LDL-C), total cholesterol, non-high-density lipoprotein (non-HDL-C), or apolipoprotein B levels)

Medication being provided by Specialty Pharmacy – Proprium Rx

Not all drugs may be covered under every Plan

If a drug is non-formulary on a Plan, documentation of medical necessity will be required.

**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. *