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COMMONWEALTH OF VIRGINIA DEPARTMENT OF MEDICAL ASSISTANCE SERVICES Service Authorization (SA) Form

Lipotropics, Other

If the following information is not complete, correct, or legible, the SA process can be delayed.

Please use one form per member.

MEMBER INFORMATION		
Last Name:	First Name:	
Medicaid ID Number:	Date of Birth:	
PRESCRIBER INFORMATION		
Last Name:	First Name:	
NPI Number:		
Phone Number:	- Fax Number:	
DRUG INFORMATION		
Is the Drug Prescribed by or in Consultation with a Cardiologists Lipidologists Endocring Drug Name/Form: Strength: Dosing Frequency: Length of Therapy: Quantity per Day:	ologists Other:	

Virginia DMAS SA Form: Lipotropics, Other

Me	ember's Last Name: Member's First Name:
CR	ITERIA
	For what indication(s) is the drug being prescribed? Check all that apply. To reduce the risk of myocardial infarction, stroke, and coronary revascularization in adults with established cardiovascular disease. As an adjunct to diet, alone or in combination with other lipid-lowering therapies (e.g., statins, ezetimibe), for treatment of adults with primary hyperlipidemia (including heterozygous familial hypercholesterolemia [HeFH]) to reduce low-density lipoprotein cholesterol (LDL-C). As an adjunct to diet and other LDL-lowering therapies (e.g., statins, ezetimibe, LDL apheresis) in patients with homozygous familial hypercholesterolemia (HoFH) who require additional lowering of LDL-C.
	The member has had prior treatment history with highest available dose or maximally-tolerated dose of high intensity statin (atorvastatin or rosuvastatin) and ezetimibe for at least three continuous months with failure to reach target LDL-C and is in one of the three groups identified by NLA (i.e., extremely high risk ASCVD members with LDL-C ≥ 70 mg/dL, very high risk atherosclerotic cardiovascular disease [ASCVD] members with LDL-C ≥ 100 mg/dL, and high-risk members with LDL-C ≥ 130 mg/dL.
	Other:
2.	Is this request for a new start or continuation of therapy? (If New Start , skip to diagnosis section.) New Start Continuation
3.	Was this drug previously authorized for this member and are they stable on the medication? (If No , skip to diagnosis section.) Yes No
4.	How long has the member been receiving treatment with these medications?
	3 to 5 months (or first renewal request after initial authorization)
	6 months or more (or second and subsequent renewal requests)
5.	For PCSK9S Leqvio®, Praluent®, or Repatha® therapy only: Has the member achieved at least a 30% reduction in LDL-C since the beginning of treatment with Leqvio®, Praluent®, or Repatha®? Action required: If Yes, please attach clinical notes and laboratory results that support reduction in LDL-C after initiation of therapy. Yes No
6.	For ATP Citrate Lyase (M4V) Nexletol® or Nexlizet™ therapy only: Has the member achieved at least a 15% to 20% reduction in LDL-C since the beginning of treatment with Nexletol® or Nexlizet™? Action required: If Yes, please attach clinical notes and laboratory results that support reduction in LDL-C after initiation of therapy. Yes No

(Form continued on next page.)

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Member's First Name:

Member's Last Name:

7.	Does the member continue to benefit from treatment as measured by either continued decrease in LDL-C levels or maintenance of optimum LDL-C levels? Action required: If Yes , please attach clinical notes and laboratory results that support continued benefit
	of therapy. Yes No
8.	
	a. Muscle symptoms resolved after discontinuation of statin; AND
	b. Muscle symptoms occurred when re-challenged at a lower dose of the same statin; AND
	c. Muscle symptoms occurred after switching to an alternative statin; AND
	d. Documentation ruling out non-statin causes of muscle symptoms (e.g., hypothyroidism, reduced renal function, reduced hepatic function, rheumatologic disorders [e.g., polymyalgia rheumatica], steroid myopathy, vitamin D deficiency, or primary muscle disease); OR
	e. The member has been diagnosed with statin-induced rhabdomyolysis
	Yes No
	If Yes to any, give details:
	AGNOSIS AND LAB VALUES FOR HOMOZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA (HOFH)
9.	Has genetic testing confirmed the presence of two mutant alleles at the LDLR, APOB, PCSK9, or LDLRAP1 gene locus?
	Action required: If Yes, please attach a copy of genetic testing result.
	Yes No
10	. Has the diagnosis of HoFH been confirmed by any of the following? Action required: Please indicate below and provide a copy of the laboratory report with LDL-C level at time of diagnosis and other documentation supporting the presence of xanthoma or family history of HoFH (e.g., chart notes, medical records).
	Untreated LDL-C > 500 mg/dL and cutaneous or tendon xanthoma before age 10 years
	Untreated LDL-C > 500 mg/dL and untreated elevated LDL-C levels consistent with heterozygous familial hypercholesterolemia in both parents
	Treated LDL-C ≥ 300 mg/dL and cutaneous or tendon xanthoma before age 10 years
	Treated LDL-C ≥ 300 mg/dL and untreated elevated LDL-C levels consistent with heterozygous familial
	hypercholesterolemia in both parents
	None of the above
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Member's First Name:

Member's Last Name:

11. Does the member have a history of clinical ASCVI ones.	D or a cardiovascular event listed below? Indicate which			
Acute coronary syndromes	Myocardial infarction			
Stable or unstable angina	Transient ischemic attack (TIA)			
Stroke of presumed atherosclerotic origin				
	 Coronary or other arterial revascularization procedure (e.g., percutaneous transluminal coronary angioplasty [PTCA], coronary artery bypass graft [CABG]) Peripheral arterial disease of presumed atherosclerotic origin Findings from a computerized tomography (CT) angiogram or catheterization consistent with clinical ASCVD 			
Peripheral arterial disease of presumed ather				
12. What is the member's pre-treatment LDL-C level	(i.e., prior to starting PCSK9 or M4V therapy)?			
mg/dL.				
Is, the member diagnosed with homozygous familial hypercholesterolemia (HoFH) and is at least 10 year of age for Repatha® or at least 18 years of age for Praluent®? Yes No				
DIAGNOSIS AND LAB VALUES FOR HETEROZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA (HEFH)				
 L4. Does the member have a definite diagnosis of heterozygous familial hypercholesterolemia (HeFH) as defined by the Dutch Lipid Clinic Network criteria (total score greater than 8)? Action required: If Yes, please provide a copy of the lab repot with LDL-C level at time of diagnosis and other documentation supporting clinical/family history and/or physical findings (e.g., chart notes, medica records). Yes No 				
15. Does the member have a definite diagnosis of He at least 10 years of age for Repatha® or at least 8 Yes No	eFH as defined by Simon Broome diagnostic criteria and is grears of age for Praluent®?			
Prescriber Signature (Required)	Date			
By signature, the physician confirms the above inform	mation is accurate and verifiable by member records.			
Please include ALL requested information; Incomple Submission of documentation does NOT guarantee co	ete forms will delay the SA process. overage by the Department of Medical Assistance Services.			
The completed form may be: FAXED TO 800-932-665 Prime Therapeutics Management LLC Attn: GV – 4201 P.O. Box 64811 St. Paul, MN 55164-0811	51 , phoned to 800-932-6648, or mailed to:			