Caterpillar Prescription Drug Benefit Phone: 877-228-7909 Fax: 800-424-7640

**Instructions:** Please fill out all applicable sections completely and legibly. Attach any additional documentation that is important for the review (e.g., chart notes or lab data, to support the authorization request). Information contained in this form is Protected Health Information under HIPAA.

JEMPER INFORMATION			U
MEMBER INFORMATION  AST NAME:		FIRST NAME:	
AST NAIVIE:		FIRST NAIVIE:	
PHONE NUMBER:		DATE OF BIRTH:	
STREET ADDRESS:			
CITY:		STATE: ZIP CODE:	
PATIENT INSURANCE ID N	UMBER:		
	EIGHT (IN/CM): WI	EIGHT (LB/KG): ALLERGIES:	
<del></del>			
YOU ARE NOT THE PATIENT OR THE PRES LLOWING LINK: PRIMETHERAPEUTICS.CO		DISCLOSURE AUTHORIZATION FORM WITH THIS REQUEST WHICH CAN BE FOUN	D AT THE
	_		
ATIENT'S AUTHORIZED RF	PRESENTATIVE (IF APPLICAR	LE):	
	TIVE'S PHONE NUMBER:		
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PRESCRIBER INFORMATIO	N		
	N	FIRST NAME:	
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Prime THERAPEUTICS\*

Revision Date: 3/15/24

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MEMBER'S LAST NAME:	MEMBER'S FIRST NAME:			
1. HAS THE PATIENT TRIED ANY OTHER	R MEDICATIONS FOR THIS CONDITION?	YES (if yes, complete below) NO		
MEDICATION/THERAPY (SPECIFY	<b>DURATION OF THERAPY</b> (SPECIFY	RESPONSE/REASON FOR		
DRUG NAME AND DOSAGE):	DATES):	FAILURE/ALLERGY:		
2. LIST DIAGNOSES:		ICD-10:		
□ Atherosclerotic Cardiovascular Disease (A	ASCVD)	100 101		
☐ Heterozygous Familial Hypercholesterole				
☐ Other diagnosis:ICD-				
3. REQUIRED CLINICAL INFORMATION: PLEASE PROVIDE ALL RELEVANT CLINICAL INFORMATION TO SUPPORT A				
PRIOR AUTHORIZATION.				
Clinical Information:				
Does the patient have any of the follow	wing:   Yes   No Please submit documen	tation		
<ul> <li>History of myocardial infarction</li> </ul>	_			
•	zation procedure more than 3 months a	go		
<ul> <li>History of unstable angina mor</li> </ul>	e than 3 months ago			
	t least one major coronary artery, as doo			
<ul> <li>Clinically significant coronary heart disease diagnosed by stress testing, as documented in a procedural report</li> </ul>				
<ul> <li>Claudication or resting limb ischemia with ankle brachial index of 0.9 or lower, as documented by an imaging</li> </ul>				
report showing at least 50% stenosis				
·				
	<ul> <li>Confirmed abdominal or thoracic aortic aneurysm</li> <li>History of lower extremity amputation</li> </ul>			
	Ischemic stroke more than 3 months ago			
	6			
•	documented in an imaging report			
	wing:   Yes   No Please submit document			
	ation in the low-density lipoprotein (LDL)	) receptor, ApoB,or PCSK9 in patient		
with untreated/ pre-treatment	nas in: (1.) patient, first degree relative,	or second degree relative with		
	cholesterol (LDL-C) >190mg/dL (age 18 ye	_		
· •	ntreated/pre-treatment LDL-C >155mg/d			
_	tient using Dutch Lipid Clinic Network di			
•	9 points (i.e.,definite FH) –(calculation w	<del>-</del>		
	_			
Is the patient currently on Statin therapy? ☐ Yes ☐ No				
If the patient is not able to use a maximum dose of a statin due to muscle symptoms, a causal relationship must be				
established between statin use and mo		iptoms, a causar relationship must be		
with the state of the sta	ujp. vaa audii ua.			



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Does the patient have evidence of pain, tenderness, stiffness, cramping, weakness, and/or fatigue and all of the
following? ☐ Yes ☐ No Please provide documentation.
Does patient have muscle symptoms that resolve after discontinuation of statin? ☐ Yes ☐ No <u>Please provide</u>
<u>documentation.</u>
Does patient have muscle symptoms occurring when re-challenged at a lower dose of the same statin? ☐ Yes ☐ No
Please provide documentation.
Did muscle symptoms occurr after switching to an alternative statin? ☐ Yes ☐ No Please provide documentation.
Has non-statin causes of muscle symptoms (e.g., hypothyroidism, reduced renal function, reduced hepatic function,
rheumatologic disorders, such as polymyalgia rheumatica, steroid myopathy, vitamin D deficiency, or primary
muscle disease) have been ruled out?   Yes   No   Please provide documentation.
Has The patient been diagnosed with rhabdomyolysis associated with statin use? ☐ Yes ☐ No Please provide
documentation.
Did the patient experience acute neuromuscular illness or dark urine and an acute elevation in creatine kinase?
Yes   No Please provide documentation.
Does the patient have a fasting LDL-C level greater than or equal to 70 mg/dL?
□ Yes □ No Please submit documentation
Does the patient have fasting triglycerides less than 500 mg/dL? ☐ Yes ☐ No Please submit documentation
Does the patient have a BMI less than 50kg/m <sup>2</sup> ? □ Yes □ No
Does the patient have a current history of renal dysfunction, nephrotic syndrome, or past history of nephritis?  □ Yes □ No
Has the patient already tolerated a trial with generic ezetemibe and/or Nexletol as indivual product(s) before reqesting this combination product? $\Box$ Yes $\Box$ No
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?
Please note: Not all drugs/diagnosis are covered on all plans. This request may be denied unless all required information is received.
<b>ATTESTATION:</b> I attest the information provided is true and accurate to the best of my knowledge. I understand that
the Health Plan, insurer, Medical Group or its designees may perform a routine audit and request the medical
information necessary to verify the accuracy of the information reported on this form.
Prescriber Signature or Electronic I.D. Verification: Date:
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### **FAX THIS FORM TO: 800-424-7640**

MAIL REQUESTS TO: Prime Therapeutics Management Prior Authorization Program
Attn: CP-4201
P.O. Box 64811
St. Paul, MN 55164-0811

Phone: 877-228-7909

