



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.

			☐ ORGENI
MEMBER INFORMATION			
LAST NAME:		FIRST NAME:	
PHONE NUMBER:		DATE OF BIRTH:	
STREET ADDRESS:			
CITY:		STATE: ZIP CODE	:
PATIENT INSURANCE ID NUI	MBER:		
MALE FEMALE HEIC IF YOU ARE NOT THE PATIENT OR THE PRESCRIFOLLOWING LINK: https://magellanrx.co	BER, YOU WILL NEED TO SUBMIT A PHI DISCLO	OSURE AUTHORIZATION FORM WITH THIS RE	QUEST WHICH CAN BE FOUND AT THE
PATIENT'S AUTHORIZED REPR			
PRESCRIBER INFORMATION			
LAST NAME:		FIRST NAME:	
PRESCRIBER SPECIALTY:		EMAIL ADDRESS:	
NPI NUMBER:		DEA NUMBER:	
PHONE NUMBER:		FAX NUMBER:	
STREET ADDRESS:			
CITY:		STATE: ZIP CODE:	
REQUESTOR (if different than prescriber):		OFFICE CONTACT PERSON:	
MEDICATION OR MEDICAL	DISPENSING INFORMATION		
MEDICATION NAME:			
DOSE/STRENGTH:	FREQUENCY:	LENGTH OF THERAPY/REFILLS:	QUANTITY:
□ NEW THERAPY □ RENEWAL IF RENEWAL: DATE THERAPY INITIATED: DURATION OF THERAPY (SPECIFIC DATES):			

Continued on next page.







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MEMBER'S LAST NAME:	MEMBER'S FIRST NAME:		
1. HAS THE PATIENT TRIED ANY OTHE	R MEDICATIONS FOR THIS CONDITION?	YES (if yes, complete below) NO	
MEDICATION/THERAPY (SPECIFY DRUG NAME AND DOSAGE):	DURATION OF THERAPY (SPECIFY DATES):	RESPONSE/REASON FOR FAILURE/ALLERGY:	
2. LIST DIAGNOSES:		ICD-10:	
 □ Clinical atherosclerotic cardiovascular □ Heterozygous familial hypercholestero □ Homozygous familial hypercholestero □ Primary hyperlipidemia 	olemia (HeFH)	165-10.	
□ Other diagnosis:	ICD-10 Code(s):		
3. REQUIRED CLINICAL INFORMATIO PRIOR AUTHORIZATION. Clinical Information:	N: PLEASE PROVIDE ALL RELEVANT CLIN	IICAL INFORMATION TO SUPPORT A	
For all diagnoses, upon initial and re	newal requests, answer the following:		
Will Repatha be used as an adjunct	to a low-fat diet and exercise? 🗆 Yes 🛭	□ No	
Is Repatha prescribed by, or in cons	ultation with, a cardiologist or endocri	nologist? 🗆 Yes 🗆 No	
Will Repatha be used in combination or Juxtapid (lomitapide)? ☐ Yes ☐ I	with another proprotein convertase su No	ubtilisin/kexin type 9 (PCSK9) inhibitor	
Will Statin therapy at a maximally to provide documentation.	lerated daily dose be continued with P	CSK9 therapy? □ Yes □ No <i>Please</i>	
Does patient have an absolute conti	raindication to statin therapy? Yes	□ No <i>Please provide documentation</i> .	
Will ezetimibe, bempedoicacid or all Please provide documentation.	oile-acid sequestrant therapy be contin	ued with PCSK9 therapy? Yes No	
Does patient have an absolute contradocumentation.	raindication to other lipid-lowering age	ents? 🗆 Yes 🗆 No Please provide	
If the patient is not able to use a max established between statin use and m	imum dose of a statin due to muscle syn ouscle symptoms such as:	nptoms, a causal relationship must be	
Does the patient have evidence of p following? Yes No Please provi	ain, tenderness, stiffness, cramping, w de documentation.	eakness, and/or fatigue <u>and all of the</u>	
Does patient have muscle symptoms documentation.	s that resolve after discontinuation of s	statin? Yes No Please provide	
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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 occur 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? 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.					
For diagnosis of Clinical atherosclerotic cardiovascular disease:					
Does patient have at least one of the following major risk factors: <i>Please provide documentation</i> .					
□ Diabetes mellitus, type 1 or 2					
□ Age 65 years or older					
□ MI or non-hemorrhagic stroke (TIAs don't qualify) in the past 6 months					
□ Current daily cigarette smoker					
□ History of more than one MI					
□ History of more than one non-hemorrhagic stroke (TIAs don't qualify)					
 History of one MI plus one non-hemorrhagic stroke (TIAs don't qualify) History of one MI plus history of symptomatic peripheral arterial disease as defined above 					
☐ History of one non-hemorrhagic stroke (TIAs don't qualify) plus history of symptomatic peripheral arterial					
disease as defined above					
IF PATIENT DOES NOT HAVE ANY OF THE ABOVE, does patient have at least 2 of the following minor risk factors					
below: Please provide documentation					
□ History of non-MI related coronary revascularization					
□ Residual coronary artery disease with >40% stenosis in at least 2 large vessels					
□ Metabolic syndrome (as defined by Alberti et al., Circulation, 2009; 120:1640-1645,					
□ Most recent HDL-C < 40 mg/dL (men) and < 50 mg/dL (women), in the absence of metabolic syndrome or in the					
presence of metabolic syndrome when 3 of its four non-HDL criteria are met (as per Alberti et al., 2009)					
□ Most recent hsCRP (high-sensitivity C-reactive protein) > 2.0 mg/L					
□ Most recent LDL-C > 130 mg/dL or non-HDL-C > 160 mg/dL					
 □ Most recent fasting LDL-C > 70 mg/dL or non-HDL-C > 100mg/dL after > 2 weeks stable lipid lowering therapy □ Most recent fasting triglycerides < 400 mg/dL 					
Is patient classified as very high risk ASCVD, defined as extensive burden of or active ASCVD, or ASCVD with extremely high burden of adverse poorly controlled cardiometabolic risk factors requiring LDL-C ≤ 70 mg/dL?					
□ Yes □ No Please provide documentation.					

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Is patient classified as high risk ASCVD, defined as less extensive ASCVD and poorly controlled cardiometabolic risk factors requiring LDL-C <100 mg/dL? Yes No Please provide documentation.				
Is patient classified as Intermediate risk ASCVD with LDL-C ≥ 130 mg/dL; with poorly controlled risk factors? □ Yes □ No Please provide documentation.				
For diagnosis of primary hyperlipidemia, please answer the following:				
Does patient have a fasting LDL-C greater than or equal to 75mg/dL? ☐ Yes ☐ No Please provide documentation.				
Does patient have a diagnosis of coronary heart disease(CHD) or is patient a risk equivalent for CHD? Please provide documentation.				
Has patient had previous background lipid-lowering therapy in which patient requires a LDL-C less than 100mg/dL? — Yes — No Please provide documentation.				
If patient does not have coronary heart disease(CHD) or is not a risk a CHD risk equivalent, has the patient had background lipid-lowering therapy requiring a LDL-C less than 130mg/dL? — Yes — No Please provide documentation.				
Does patient have a triglyceride level less than or equal to 400mg/dL? Yes No Please provide documentation.				
Is patient NYHA class II, III or IV? □ Yes □ No Please provide documentation.				
Is patient's last known left ventricular ejection fracture less than 30%? ☐ Yes ☐ No				
Is patient a Type I diabetic? ☐ Yes ☐ No				
Is patient a poorly controlled Type II diabetic with a HgA1c greater than or equal to 7%? ☐ Yes ☐ No				
Does patient have uncontrolled hypertension with a blood pressure greater than or equal to 140/90mmHg?□ Yes □ No				
SEE BELOW FOR ADDITIONAL SECTIONS FOR HEFH AND HoFH:				
For heterozygous familial hypercholesterolemia (HeFH), also answer the following: Has there been genetic confirmation of the diagnosis through a mutation identified in the LDL receptor, ApoB or PCSK9? Yes No Please provide documentation				
If yes, does patient have an untreated/pre-treatment LDL-C greater than 190 mg/dL?				
Is there documented evidence of tendinous xanthomas in the patient and/or first-degree relative, and/or second-degree relative? □ Yes □ No Please provide documentation				
If Yes, is the individual with tendinous xanthomas a first- or second-degree relative less than 18 years of age with an untreated/pre-treatment LDL-C greater than 155 mg/mL? □ Yes □ No Please provide documentation				

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If Yes, is the individual with tendinous xanthomas 18 years of age or older with an untreated/pre-treatment LDL-C greater than 190 mg/mL? ☐ Yes ☐ No Please provide documentation
Was the patient assessed with the Dutch Lipid Clinic Network diagnostic criteria and found to have a cumulative
score greater than or equal to 9 points (i.e., definite FH)?
Does the patient's fasting LDL-Cvalue within the last 30 days while on a maximally tolerated lipid-lowering regimen equal 100mg/dL or greater? No Please provide documentation
Is the fasting triglyceride level for this patient greater than 400 mg/dL? — Yes — No Please provide documentation
For homozygous familial hypercholesterolemia (HoFH), also answer the following:
Has there been genetic confirmation of the diagnosis through two mutant alleles identified in the LDL receptor, ApoB, PCSK9 or LDL receptor adaptor protein 1 (LDLRAP1 or ARH)? ☐ Yes ☐ No Please provide documentation
Does the patient have a untreated/pre-treatment LDL-C greater than 500 mg/dL? — Yes — No Please provide documentation
Did the patient have a cutaneous or tendinous xanthoma before the age of ten? — Yes — No — Please provide documentation
Do both of the patient's parents have evidence of heterozygous familial hypercholesterolemia? — Yes — No — No
Has the patient had cellular testing performed which demonstrated a reduced LDL receptor activity in fibroblasts/lymphocytes equaling 20% or less of the normal activity? ☐ Yes ☐ No Please provide documentation
Does the patient's fasting LDL-Cvalue within the last 30 days while on a maximally tolerated lipid-lowering regimen equal 130mg/dL or greater? \[\text{Yes} \text{No} \text{Please provide documentation} \]
Is the fasting triglyceride level for this patient greater than 400 mg/dL? ☐ Yes ☐ No Please provide documentation
REAUTHORIZATIONS:
If this is a reauthorization request, answer the following questions:
$Have\ medical\ records\ (e.g.,\ laboratory\ values)\ been\ submitted\ that\ document\ a\ sustained\ reduction\ in\ LDL-C\ levels$
from pre-treatment baseline (i.e., prior to PCSK9 therapy) while on PCSK9 therapy? ¬ Yes ¬ No

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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.				
ATTESTATION: 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: Magellan Rx Management Prior Authorization Program

Attn: CP – 4201 P.O. Box 64811 St. Paul, MN 55164-0811

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