## AvMed

## 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 1-305-671-0200</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>

Drug Re	equested: (select drug below)	
□ Nexl	etol <sup>™</sup> (bempedoic acid)	□ Nexlizet <sup>™</sup> (bempedoic acid/ezetimibe)
MEMB	BER & PRESCRIBER INFORMAT	ION: Authorization may be delayed if incomplete.
Member 1	Name:	
	AvMed #:	
Prescribe	r Name:	
Prescribe	r Signature:	Date:
Office Co	ntact Name:	
Phone Number:		
DEA OR	NPI #:	
DRUG	<b>INFORMATION:</b> Authorization may 1	be delayed if incomplete
Dosing So	chedule:	Length of Therapy:
Diagnosis	:	ICD Code, if applicable:
Weight:		Date:
support e		apply. All criteria must be met for approval. To ng lab results, diagnostics, and/or chart notes, must be
Initial A	Authorization: 6 months	
□ Mı	ust be prescribed or in consultation with one	of the following:
	Cardiologist	
	Endocrinologist	
u	Lipid Specialist	

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<ul><li>A</li><li>H</li><li>O</li></ul>	CD-10 Code(s) plus description:
ca	Atherosclerotic Cardiovascular Disease – Select if the member has Atherosclerotic ardiovascular disease (ASCVD) confirmed by the following: (Please note: Chart documentation is equired to be submitted along with this request form.)
	Acute coronary syndrome
	History of myocardial infarction
	Stable or unstable angina
	Coronary artery disease
	Coronary or other arterial revascularization (e.g., percutaneous coronary intervention (PCI), angioplasty coronary stent procedure or coronary bypass graft (CABG) surgery)
	Stroke
	Transient ischemic attack
	Peripheral arterial disease presumed to be of atherosclerotic origin
	<u>OR</u>
Н	<b>leterozygous Familial Hypercholesterolemia (HeFH)</b> – Select if the member has eterozygous familial hypercholesterolemia (HeFH) confirmed by the following: <b>(Please note: Chart ocumentation is required to be submitted along with this request form.)</b>
	Untreated/pre-treatment LDL cholesterol (LDL-C) $\geq$ 190mg/dL in an adult or $\geq$ 155mg/dL in a child less than 16 years of age
	AND (ONE OF THE FOLLOWING)
	Family history of myocardial infarction in first-degree relative less than 60 years of age
	Family history of myocardial infarction in second-degree relative less than 50 years of age
	Family history of familial hypercholesterolemia in first-or second degree relative
	Submission of medical records (e.g., chart notes, laboratory values) documenting LDL-C > 190mg/dL
	in first or second degree relative.ransient ischemic attack
	OR (ONE OF THE FOLLOWING)

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Genetic confirmation of functional mutation in the LDL receptor, Apo-B, or PCSK9 gene adaptor protein 1 (i.e., LDLRAP1 or ARH)				
Tendinous xanthomata				
Arcus cornealius before age 45				
AND				
lease confirm ALL the follow ocumentation is required to be sub	0	and/or HeFH: (Please note: Chart this request form.)		
	less the addition of e	statin 40-80mg daily, rosuvastatin 20-40mg daily ezetimibe is contraindicated) for > 12 continuous		
<u>OR</u>				
	by and is on maximally tolerated statin therapy ezetimibe is contraindicated) for > 12 continuous			
Statin:	_ Strength:	Date started:		
AND (ONE OF	THE FOLLOW	ING)		
LDL-C remains greater than or equal to 70 mg/dL with ASCVD				
LDL-C remains greater than or equal to 100 mg/dL without ASCVD				
**Please document: the LDL level delayed)**	els below (Labs <mark>M</mark>	IST be attached or authorization will be		
LDL baseline:	LI	OL post therapy:		
<u>OR</u>				
	fferent statins (i.e., t	d by <u>ONE</u> of the following intolerable and rial of at least 14 days of each) (documentation		
☐ Myalgia (muscle symptoms without CK elevations) <u>OR</u>				
☐ Myositis (muscle symptoms wi	th CK elevations < 1	10 times upper limit of normal)		
<u>AND</u>				
Reinitiating of statin therapy at low attempted and failed (documentation)		requency of administration must have been ance MUST be provided)		
**Please document statin therapy below; pharmacy claims will be verified**				
Statin:	Strength:	Date started:		
Statin:	Strength:	Date started:		
<u>OR</u>				

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	experienced rhabd	eled contraindication to ALL statins as documented in medical records and/or has lomyolysis or muscle symptoms with statin treatment with CK elevations > 10 times mal (documentation of labeled contraindication to ALL statins must be provided)		
	$\underline{\mathbf{A}}\mathbf{N}$	<u>ND</u>		
	authorization) and	<u>90-Day</u> trial of a PCSK9 inhibitor (i.e., Repatha <sup>®</sup> or Praluent <sup>®</sup> - require prior failed to reach LDL target goal (documentation of PCSK9 inhibitor failure, bs after 90 days of therapy, MUST be provided)		
	<u>OI</u>	<u>R</u>		
	Member has had a life-threatening adverse reaction to a PCSK9 inhibitors (i.e., Repatha® or Praluent® – required prior authorization) (documentation of life-threatening adverse reaction MUST be provided)			
**Pl	ease note: Conco	omitant therapy with PCSK9 inhibitors will not be approved**		
supp		<b>2 months.</b> Check below all that apply. All criteria must be met for approval. To ed, all documentation, including lab results, diagnostics, and/or chart notes, must be be denied.		
	Documentation of	positive clinical response to therapy (e.g., reduction in LDL-C levels)		
	AN	ND		
	Member continues tolerated dose	s to receive other lipid-lowering therapy (e.g., statin, ezetimibe) at a maximally		
	OI	₹		
	Member has a doc	rumented inability to take other lipid-lowering therapy (e.g., statins, ezetimibe)		
	**Please docume delayed)**	nt: the LDL levels below (Labs MUST be attached or authorization will be		
	LDL baseline:	LDL post-Nexletol/Nexlizet:		

Medication being provided by a 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 phi rmacy paid claims or submitted chart notes.\*