



Repatha

HMSA - Prior Authorization Request

CVS Caremark administers the prescription benefit plan for the patient identified. This patient's benefit plan requires prior authorization for certain medications in order for the drug to be covered. To make an appropriate determination, providing the most accurate diagnosis for the use of the prescribed medication is necessary. **Please respond below and fax this form to CVS Caremark toll-free at 1-866-237-5512.** If you have questions regarding the prior authorization, please contact CVS Caremark at **1-808-254-4414**. For inquiries or questions related to the patient's eligibility, drug copay or medication delivery; please contact the Specialty Customer Care Team: CaremarkConnect® 1-800-237-2767.

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Patient's Name: _____ **Date:** _____
Patient's ID: _____ **Patient's Date of Birth:** _____
Patient's Phone Number: _____
Physician's Name: _____
Specialty: _____ **NPI#:** _____
Physician Office Telephone: _____ **Physician Office Fax:** _____

Approvals may be subject to dosing limits in accordance with FDA-approved labeling, accepted compendia, and/or evidence-based practice guidelines.

Additional Demographic Information:

Patient Weight: _____ *kg*
Patient Height: _____ *ft* _____ *inches*

Indicate where the drug is being dispensed:

- Office Outpatient Hospital Ambulatory Surgical Inpatient Hospital
- Off Campus Outpatient Hospital Urgent Care Emergency Room Birthing Center
- Military Facility Skilled Nursing Facility Nursing Facility Hospice
- Inpatient Psychiatric Psychiatric Residential Treatment End Stage Renal Facility
- Psychiatric Facility Pharmacy Other

Indicate where the drug is being administered:

- Ambulatory surgical Home Inpatient hospital Office
- Outpatient Hospital Pharmacy

What is the ICD-10 code? _____

Send completed form to: CVS Caremark Specialty Programs. Fax: 1-866-237-5512

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Criteria Questions:

1. Does the patient have a history of clinical atherosclerotic cardiovascular disease (ASCVD)?

Yes, *Continue to #2*

No, *Continue to #3*

2. Is this a request for continuation of therapy with a PCSK9 inhibitor?

Yes, *Continue to #100*

No, *Continue to #200*

3. What is the diagnosis?

Primary hyperlipidemia, *Continue to #4*

Heterozygous familial hypercholesterolemia (HeFH), *Continue to #4*

Homozygous familial hypercholesterolemia (HoFH), *Continue to #4*

Other, *Continue to #4*

4. Is this a request for continuation of therapy with a PCSK9 inhibitor?

Yes, *Continue to #100*

No, *Continue to #300*

Continuation

100. Has the patient achieved or maintained an LDL-C reduction (e.g., LDL-C is now at goal, robust lowering in LDL-C) as a result of PCSK9 inhibitor therapy?

Yes, *No Further Questions*

No, *No Further Questions*

ASCVD

200. Which of the manifestations of clinical atherosclerotic cardiovascular disease (ASCVD) has the patient experienced?

Acute coronary syndrome(s), *Continue to #201*

Myocardial infarction, *Continue to #201*

Stable or unstable angina, *Continue to #201*

Coronary or other arterial revascularization procedure (e.g., percutaneous coronary intervention [PCI], coronary artery bypass graft [CABG] surgery), *Continue to #201*

Stroke of presumed atherosclerotic origin, *Continue to #201*

Transient ischemic attack (TIA), *Continue to #201*

Non-cardiac peripheral arterial disease (PAD) of presumed atherosclerotic origin (e.g., carotid artery stenosis, lower extremity PAD), *Continue to #201*

Other, *Continue to #201*

201. What is the current LDL-C level?

_____ mg/dL, *Continue to #600*

Unknown, *Continue to #600*

Primary Hyperlipidemia/HoFH/HeFH

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300. What is the patient's untreated (before any lipid-lowering therapy) LDL-C level?

_____ mg/dL, *If greater than or equal to 190mg/dL continue to #303, If less than 190mg/dL continue to #301*

Unknown, *Continue to #303*

301. Does the patient have a diagnosis of heterozygous familial hypercholesterolemia or homozygous familial hypercholesterolemia?

Yes, *Continue to #302*

No, *Continue to #302*

302. Is the patient less than 18 years of age?

Yes, *Continue to #303*

No, *Continue to #303*

303. Are there any secondary causes that could explain the elevated untreated LDL-C?

Yes, *Continue to #304*

No, *Continue to #304*

304. What is the current LDL-C level?

_____ mg/dL, *Continue to #600*

Unknown, *Continue to #600*

Previous Lipid Lowering Therapy

600. Is the patient receiving a high-intensity statin dose daily, such as rosuvastatin (Crestor) 20 mg daily or atorvastatin (Lipitor) 40 mg daily?

Yes, *Continue to #601*

No, *Continue to #603*

601. Has the patient received this dose for at least 3 months?

Yes, *Continue to #602*

No, *Continue to #603*

602. Has the patient received the high-intensity statin dose for at least 3 months in combination with ezetimibe?

Yes, *No Further Questions*

No, *No Further Questions*

603. Was the patient unable to tolerate a high-intensity statin due to adverse effects?

Yes, *Continue to #604*

No, *Continue to #650*

604. Is the patient receiving a moderate-intensity statin dose daily, such as atorvastatin (Lipitor) 20 mg daily or equivalent?

Yes, *Continue to #605*

No, *Continue to #650*

605. Has the patient received this dose for at least 3 months?

Yes, *Continue to #606*

No, *Continue to #650*

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606. Has the patient received the moderate-intensity statin dose for at least 3 months in combination with ezetimibe?

Yes, *No Further Questions*

No, *No Further Questions*

Intolerance and Contraindications

650. Did the patient score a 7 or higher on the Statin-Associated Muscle Symptom Clinical Index (SAMS-CI)?

Yes, *No Further Questions*

No, *Continue to #651*

651. Did the patient experience a statin-associated increase in creatine kinase level of greater than or equal to 10 times the upper limit of normal (ULN) during previous treatment with a statin?

Yes, *No Further Questions*

No, *Continue to #652*

652. Does the patient have any of the following contraindications to statins?

Active liver disease, including unexplained persistent elevations in hepatic transaminase levels (e.g., ALT greater than or equal to 3 times upper limit of normal), *No Further Questions*

Currently pregnant, *No Further Questions*

Planning pregnancy, *No Further Questions*

Breastfeeding, *No Further Questions*

None of the above, *No Further Questions*

I attest that this information is accurate and true, and that documentation supporting this information is available for review if requested by CVS Caremark or the benefit plan sponsor.

X _____

Prescriber or Authorized Signature

Date (mm/dd/yy)

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Statin-Associated Muscle Symptom Clinical Index (SAMS-CI)

Instructions:

- Use with patients who have had muscle symptoms that were **new or increased** after starting a statin regimen.
- A **statin regimen** includes any statin at any dose or frequency, including a statin the patient has used previously, at the same or a different dose.
- **Muscle symptoms** may include aches, cramps, heaviness, discomfort, weakness, or stiffness.
- Interpret overall score in light of **other possible causes** of the muscle symptoms, such as:

Recent physical exertion	Hypothyroidism	Concurrent illness
Changes in exercise patterns	Drug interaction with statin	Underlying muscle disease
- **See reverse** for Frequently Asked Questions

How many statin regimens has the patient had that involved new or increased muscle symptoms?

One

Two or more

Complete the questions on the left side of this page.

Complete the questions on the right side of this page.

Regarding this statin regimen:

A. Location and pattern of muscle symptoms

(If more than one category applies, record the highest number.)

Enter score:

Symmetric, hip flexors or thighs	3	<input style="width: 40px; height: 30px;" type="text"/>
Symmetric, calves	2	
Symmetric, proximal upper extremity	2	
Asymmetric, intermittent, or not specific to any area	1	

B. Timing of muscle symptom onset in relation to starting statin regimen

<4 weeks	3	<input style="width: 40px; height: 30px;" type="text"/>
4–12 weeks	2	
>12 weeks	1	

C. Timing of muscle symptom improvement after withdrawal of statin (If patient is still taking statin, stop regimen and monitor symptoms.)

<2 weeks	2	<input style="width: 40px; height: 30px;" type="text"/>
2–4 weeks	1	
No improvement after 4 weeks	0	

Rechallenge the patient with a statin regimen, (even if same statin compound or regimen as above) then complete final question:

D. Timing of recurrence of similar muscle symptoms in relation to starting second regimen

<4 weeks	3	<input style="width: 40px; height: 30px;" type="text"/>
4–12 weeks	1	
>12 weeks or similar symptoms did not reoccur	0	

Total:

All four scores above must be entered before totaling

Regarding the statin regimen before the most recent regimen:

A. Location and pattern of muscle symptoms (If more than one category applies, record the highest number.)

Enter score:

Symmetric, hip flexors or thighs	3	<input style="width: 40px; height: 30px;" type="text"/>
Symmetric, calves	2	
Symmetric, proximal upper extremity	2	
Asymmetric, intermittent, or not specific to any area	1	

B. Timing of muscle symptom onset in relation to starting statin regimen

<4 weeks	3	<input style="width: 40px; height: 30px;" type="text"/>
4–12 weeks	2	
>12 weeks	1	

C. Timing of muscle symptom improvement after withdrawal of statin

<2 weeks	2	<input style="width: 40px; height: 30px;" type="text"/>
2–4 weeks	1	
No improvement after 4 weeks	0	

Regarding the most recent statin regimen:

(even if same statin compound as above)

D. Timing of recurrence of similar muscle symptoms in relation to starting regimen

<4 weeks	3	<input style="width: 40px; height: 30px;" type="text"/>
4–12 weeks	1	
>12 weeks or similar symptoms did not reoccur	0	

Total:

All four scores above must be entered before totaling

	Total score:	2–6	7–8	9–11
Interpretation	Likelihood that the patient's muscle symptoms are due to statin use:	Unlikely	Possible	Probable

10 Oct 2016. Based on Rosenson et al. An assessment by the Statin Muscle Safety Task Force: 2014 update. *J Clin Lipidol.* 2014 May–Jun;8(3 Suppl):S58–71.

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