

Pharmacy Section

Member Name: _____ Date of Birth: _____ Member ID#: _____
 Pharmacy NPI: _____ Pharmacy Phone: _____ Pharmacy Fax: _____
 Pharmacy Name: _____ Pharmacist Name: _____
 Prescriber NPI: _____ Prescriber Name: _____ Specialty: _____
 Prescriber Phone: _____ Prescriber Fax: _____ Drug Name/Strength: _____
 NDC: _____ Regimen: _____ Fill Date: _____ Quantity: _____ Day Supply: _____
 Has member been trained on proper administration and storage of this medication? Yes _____ No _____
 Pharmacist Signature: _____ Date: _____

Prescriber Section

Page 1 of 2—Please complete and return all pages. Failure to complete all pages will result in processing delays.
All information must be provided and SoonerCare may verify through further requested documentation. The member's prescription claim history will be reviewed prior to approval.

For Initial Authorization (Initial approval will be for the duration of 3 months):

1. Please indicate member's diagnosis:

- ☐ Heterozygous familial hypercholesterolemia (HeFH) confirmed by: **(check all that apply)**
- ☐ Documented functional mutation(s) in low-density lipoprotein (LDL) receptor alleles or alleles known to affect LDL receptor functionality via genetic testing **(results of genetic testing must be submitted)**
 - ☐ Pre-treatment total cholesterol >290mg/dL or LDL-cholesterol (LDL-C) >190mg/dL
 - ☐ History of tendon xanthomas in either the member, first degree relative, or second degree relative
 - ☐ Dutch Lipid Clinic Network Criteria score of >8
- ☐ Homozygous familial hypercholesterolemia (HoFH) confirmed by 1 or more of the following:
- ☐ Documented functional mutation(s) in both LDL receptor alleles or alleles known to affect LDL receptor functionality via genetic testing **(results of genetic testing must be submitted)**
 - ☐ Untreated LDL-C >500mg/dL and at least 1 of the following:
 - ☐ Documented evidence of definite HeFH in both parents
 - ☐ Presence of tendinous/cutaneous xanthoma prior to 10 years of age
- ☐ To reduce the risk of myocardial infarction, stroke, coronary revascularization, and/or unstable angina requiring hospitalization in adults with established cardiovascular disease (CVD). Please provide supporting diagnoses/conditions and dates of occurrence signifying established CVD:
- Diagnosis/condition: _____ Date of occurrence: _____
- Diagnosis/condition: _____ Date of occurrence: _____
- ☐ Primary hyperlipidemia

2. How will this medication be used? ☐ Monotherapy ☐ Adjunct to statin therapy, diet, and exercise

3. Please specify the member's current statin therapy:

- a) Medication/strength: _____ Dosing regimen: _____ Duration of treatment: _____
- b) Has member been adherent to high-dose statin therapy for at least 12 continuous weeks? Yes _____ No _____
- c) If yes, please provide member's LDL-C level following 12 weeks of statin therapy: _____
SoonerCare claims analysis will be conducted to verify adherence.

Page 1 of 2

Fax completed prior authorization request form to 888-601-8461 or submit Electronic Prior Authorization through CoverMyMeds® or SureScripts. All requested data must be provided. Incomplete forms or forms without the chart notes will be returned. Pharmacy Coverage Guidelines are available at AetnaBetterHealth.com/Oklahoma.

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Member Name: _____ Date of Birth: _____ Member ID#: _____

Prescriber Section***Page 2 of 2—Please complete and return all pages. Failure to complete all pages will result in processing delays.*
For Initial Authorization, Continued:**

4. If the member has **not** been adherent to high-dose statin therapy for at least 12 continuous weeks, is the member intolerant to statin therapy? Yes ____ No ____
- a) If yes, please indicate 1 of the following:
- ☐ Rhabdomyolysis - creatine kinase (CK) labs verifying this diagnosis must be provided.
 - ☐ An FDA labeled contraindication to all statins. Provide contraindication: _____
 - ☐ Documented intolerance to at least 2 different statins at lower doses or at intermittent dosing:
Please provide all of the following:
- 1) Medication/strength: _____ Dosing regimen: _____
Duration of treatment: _____ Reason for discontinuation: _____
- 2) Medication/strength: _____ Dosing regimen: _____
Duration of treatment: _____ Reason for discontinuation: _____
5. Has the member had a recent trial of a statin with ezetimibe? Yes ____ No ____
- a) If yes, please provide statin tried with ezetimibe: _____ trial dates: _____
6. If the member is intolerant to statin therapy, has the member had a recent trial of ezetimibe alone? Yes ____ No ____
- a) If yes, please provide ezetimibe trial dates: _____
7. Please provide member's LDL-C level following ezetimibe therapy with statin therapy or without statin therapy: _____
8. If ezetimibe has not been tried either with or without a statin, please provide a patient-specific, clinically significant reason why ezetimibe is not appropriate for the member: _____
9. Member's baseline LDL-C: _____ Current LDL-C: _____ Goal LDL-C: _____
10. Has the member been counseled on proper administration and storage of PCSK9 therapy? Yes ____ No ____

For Continued Authorization:

1. Has member been compliant with PCSK9 Inhibitor treatment? Yes ____ No ____
2. Has PCSK9 Inhibitor treatment been effective for this member? Yes ____ No ____
3. Please provide a recent LDL-C level for this member: _____ Date taken: _____

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Prescriber Signature: _____ **Date:** _____*By signature, the physician confirms the criteria information above is accurate and verifiable in patient records. Please do not send in chart notes. Specific information will be requested if necessary. Failure to complete this form in full will result in processing delays.*

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