US Family Health Plan Prior Authorization Request Form for evolocumab (Repatha)

To be completed and signed by the prescriber. To be used only for prescriptions which are to be filled through the Department of Defense (DoD) US Family Health Plan Pharmacy Program. US Family Health Plan is a TRICARE contractor for DoD.

The completed form may be faxed to 855-273-5735

OR

The patient may attach the completed form to the prescription and mail it to:

Attn: Pharmacy, 77 Warren St, Brighton, MA 02135

QUESTIONS? Call 1-877-880-7007

Medical documentation may be required. Failure to provide could result in denial. Step Please complete patient and physician information (please print): 1 Patient Name: Physician Name: Address: Address: Sponsor ID# Phone #: Secure Fax #: Date of Birth: Step Please complete the clinical assessment: 2 1. What is the diagnosis or ☐ Homozygous familial hypercholesterolemia (HoFH) - Proceed to question 2 indication? ☐ Heterozygous familial hypercholesterolemia (HeFH) - Proceed to guestion 3 ☐ Clinical atherosclerotic cardiovascular disease (ASCVD) - Proceed to question 4 ☐ Other - STOP - Coverage not approved ☐ Yes Is the patient greater than or equal to 10 years of age? □ No STOP Proceed to question 5 Coverage not approved ☐ Yes □ No 3. Is the patient greater than or equal to 10 years of age? STOP Proceed to question 6 Coverage not approved ☐ Yes □ No Is the patient greater than or equal to 18 years of age? **STOP** Proceed to question 6 Coverage not approved ☐ Yes □ No Is the patient receiving other LDL-lowering therapies (for example, statin, ezetimibe, LDL apheresis), and requires **STOP** Proceed to question 7 additional lowering of LDL cholesterol? Coverage not approved ☐ Yes □ No Will the patient be on concurrent statin therapy at a maximal tolerated dose while on the requested Proceed to question 13 Proceed to question 9 medication? ☐ Yes □ No 7. Is the patient pregnant or breastfeeding? **STOP** Proceed to question 8 Coverage not approved

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9.	Is the prescribed dosage documented as 140 mg every 2 weeks, OR 420 mg every 4 weeks? Note: Only patients with homozygous familial hypercholesterolemia (HoFH) will be allowed to use 3 of the 140 mg syringes to make the 420 mg dose. Has the patient experienced intolerable and persistent (for		☐ Yes Sign and date below ☐ Yes	□ No STOP Coverage not approved □ No
	longer than 2 weeks) muscle symptoms (muscle pain, weakness, cramps) while on statin therapy?		Proceed to question 10	Proceed to question 12
10.	Has the patient undergone at least 2 trials of statin re- challenges with reappearance of muscle symptoms? NOTE: that is, the patient has had 2 trials of statins with muscle symptoms.		☐ Yes Proceed to question 13	☐ No Proceed to question 11
11.	Has the patient had a creatine kinase (CK) level greater than 10 times the upper limit of normal OR rhabdomyolysis with CK greater than 10,000 international units per liter (IU/L) that is unrelated to statin use?		☐ Yes Proceed to question 13	☐ No Proceed to question 12
12.	Does the patient have a contraindication to the use of a statin? NOTE: Please select the option that best applies to this patient's condition.		□ Active Liver Disease (including unexplained persistent elevations in hepatic transaminase levels) - Proceed to question 13 □ Hypersensitivity - Proceed to question 13 □ Pregnancy - Proceed to question 13 □ Nursing mothers - Proceed to question 13 □ None of the above - STOP - Coverage not approved	
13.	3. Is the patient pregnant or breastfeeding?		☐ Yes STOP Coverage not approved	□ No Proceed to question 14
14.	. What is the indication or diagnosis?	☐ Clinical atherosclerotic cardiovascular ☐ Heterozygous familial hypercholestero	,	•
15.	Is the prescribed dosage documented as 140 mg every 2 weeks, OR 420 mg every 4 weeks?		□ Yes Sign and date below	□ No STOP Coverage not approved
16.	Is the patient at very high risk for future ASCVD events? NOTE: patients at very high risk for future ASCVD events include those with a history of multiple major ASCVD events or 1 major ASCVD event and multiple high risk conditions. Refer to the 2022 ACC Expert Consensus Decision Pathway on the role of nonstatin therapies for LDL-cholesterol lowering in the management of ASCVD for more information.		☐ Yes Proceed to question 17	☐ No Proceed to question 18
17.	Does the patient have an LDL level greater than 55 mg/dL despite statin at maximal tolerated doses?		☐ Yes Proceed to question 19	☐ No STOP Coverage not approved
18.	Does the patient have an LDL level greater than 70 mg/dL despite statin at maximal tolerated doses?		☐ Yes Proceed to question 19	☐ No STOP Coverage not approved
19.	Has the patient tried EITHER atorvastatin (Lipitor) at a dose of 40 mg to 80 mg OR rosuvastatin (Crestor) at a dose of 20 mg to 40 mg for at least 4 to 6 weeks each?		☐ Yes Proceed to question 22	☐ No Proceed to question 20
20.		statin at a maximally tolerated ezetimibe (Zetia) for at least 4 to	☐ Yes Proceed to question 22	☐ No Proceed to question 21

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	21. Has the patient tried ezetimibe (Zetia) as monotherapy (alone) for at least 4 to 6 weeks?	☐ Yes Proceed to question 22	□ No STOP Coverage not approved		
	22. Is the prescribed dosage documented as 140 mg every 2 weeks, OR 420 mg every 4 weeks?	☐ Yes Sign and date below	☐ No STOP Coverage not approved		
Step 3	I certify the above is true to the best of my knowledge. Please sign and date:				
	Prescriber Signature	Date			
			[30 Aug 2023]		