

RepathaReady® Intake Form

Please see Repatha® (evolocumab) Indications and Important Safety Information on page 2.

Fax with copies of insurance card(s), front and back, and appropriate information from patient's medical charts to: 1-855-REPATHA (1-855-737-2842).



Patient Information

Patient Name*: _____	Preferred Phone*: () _____
Street Address*: _____	Email Address: _____
City*: _____ State*: _____ Zip*: _____	Date of Birth*: _____ Gender: <input type="checkbox"/> Male <input type="checkbox"/> Female

Pharmacy Insurance Information **Primary/Secondary Medical Insurance Information** (ONLY if Pharmacy Insurance Information is not available)

<input type="checkbox"/> Attach a copy of pharmacy card, front and back, AND provide: Pharmacy Insurance ID #: _____ Pharmacy Insurance Telephone*: () _____	<input type="checkbox"/> Attach a copy of insurance card, front and back, AND provide: Name of Insurer*: _____ Insurer Telephone: () _____ Group Number: _____ Policy Number*: _____
---	--

Prescriber Information

Office Contact: _____	Office Street Address*: _____
Email Address: _____	City*: _____ State*: _____ Zip*: _____
Prescriber Name*: _____	Telephone*: () _____
Specialty: _____	Fax: () _____
Office Name*: _____	Prescriber NPI #: _____ Tax ID #: _____

Prescription Information: Repatha®

FORMULATION AND DIRECTIONS (Choose one): <input type="checkbox"/> Inject 140-mg/mL subcutaneously using a SureClick® autoinjector every two (2) weeks† <input type="checkbox"/> Administer 420-mg/3.5 mL subcutaneously using a Pushtronex® system (on-body infuser with prefilled cartridge) once (1) monthly‡	DISPENSED QUANTITY FOR: <input type="checkbox"/> 28 days <input type="checkbox"/> 84 days <input type="checkbox"/> _____ days <input type="checkbox"/> 30 days <input type="checkbox"/> 90 days <input type="checkbox"/> _____ days	REFILLS: _____ refills _____ refills
---	---	--

I authorize Amgen and its agents to act on my behalf for the limited purposes of transmitting this prescription to the appropriate pharmacy consistent with the patient's benefit plan.

Prescriber Signature (no stamps) _____ _____ Date: _____
Dispense as written **Substitution allowed**

Patient Information

Please provide one **primary** and one **secondary** established CVD ICD-10-CM code*[§]:

Primary Codes:	Secondary Codes:	<input type="checkbox"/> I63.____ Cerebral Infarction	<input type="checkbox"/> I73.9 Peripheral Vascular Disease, Unspecified
<input type="checkbox"/> E78.00 Pure Hypercholesterolemia, unspecified	<input type="checkbox"/> I20.0 Unstable Angina	<input type="checkbox"/> I65.____ Occlusion and Stenosis of Cerebral Arteries, Extracranial	<input type="checkbox"/> G45.9 Transient Cerebral Ischemic Attack, Unspecified
<input type="checkbox"/> E78.01 Familial Hypercholesterolemia	<input type="checkbox"/> I20.9 Angina Pectoris, Unspecified	<input type="checkbox"/> I66.____ Occlusion and Stenosis of Cerebral Arteries, Intracranial	<input type="checkbox"/> G46.____ Vascular Syndromes
<input type="checkbox"/> E78.2 Mixed Hyperlipidemia	<input type="checkbox"/> I21.____ Acute Myocardial Infarction	<input type="checkbox"/> I67.____ Other Cerebrovascular Diseases	<input type="checkbox"/> Z83.42 Family history of familial hypercholesterolemia
<input type="checkbox"/> E78.4 Other Hyperlipidemia	<input type="checkbox"/> I22.____ Subsequent Myocardial Infarction	<input type="checkbox"/> I70.____ Atherosclerosis	<input type="checkbox"/> Other (specify ICD-10-CM): _____
<input type="checkbox"/> E78.5 Hyperlipidemia, Unspecified	<input type="checkbox"/> I25.____ Chronic Ischemic Heart Disease		
<input type="checkbox"/> Other (specify ICD-10-CM): _____			

Lipid-lowering Treatment History

LDL-C on treatment: _____ Date of Lab: [\ \] LDL-C on Repatha® (if applicable) _____ Date of Lab: [\ \]

Atorvastatin (Lipitor®) 10mg 20mg 40mg 80mg Date of Initiation [\ \] Treatment End Date [\ \]
 Achieved maximum tolerated statin dose? Y N

Rosuvastatin (Crestor®) 5mg 10mg 20mg 40mg Date of Initiation [\ \] Treatment End Date [\ \]
 Achieved maximum tolerated statin dose? Y N

Simvastatin (Zocor®) 5mg 10mg 20mg 40mg Date of Initiation [\ \] Treatment End Date [\ \]
 Achieved maximum tolerated statin dose? Y N

Other statin/lipid-lowering medication(s): _____ Date of Initiation [\ \] Treatment End Date [\ \]
 Achieved maximum tolerated statin dose? Y N

Ezetimibe (Zetia®) Date of Initiation [\ \] Treatment End Date [\ \] Added to statin? Y N Indicate statin: _____ Dose (mg) []

Repatha® was prescribed by, or in consultation with, a cardiologist, an endocrinologist, and/or a physician who focuses on the management of cardiovascular disease and/or lipid disorders.

Has the patient failed on or do they have contraindications to any of the above therapies? _____

Other pertinent medical history or drug therapy including allergies: _____

History of atherosclerotic cardiovascular disease (established CVD or FH): _____

* Required for processing.
 † Dosage for primary hyperlipidemia indication, including HeFH indication. The needle cover of the single-use prefilled autoinjector contains dry natural rubber (a derivative of latex) that may cause allergic reactions in individuals sensitive to latex.
 ‡ Dosage for primary hyperlipidemia, including HeFH, or for HoFH indications. The single-use on-body infuser with prefilled cartridge is not made with natural rubber latex.
 § The sample diagnosis codes are informational and not intended to be directive or a guarantee of reimbursement, and include potential codes that would include FDA-approved indications for Repatha®. Other codes may be more appropriate given internal system guidelines, payer requirements, practice patterns, and the services rendered.
 By signing above, you represent that your patient is aware of the disclosure of their personal health information to Amgen and its agents for Amgen's patient support services, including reimbursement and verification services and the services provided by field reimbursement professionals in your office, as part of the patient's treatment with this product and that you have obtained appropriate patient authorizations as needed.
 Repatha® and RepathaReady® are trademarks of Amgen Inc. All other marks used herein are the property of their respective owners. RepathaReady®: Personalized support services for patients and providers.





INDICATIONS AND IMPORTANT SAFETY INFORMATION

Prevention of Cardiovascular Events: In adults with established cardiovascular disease, Repatha® is indicated to reduce the risk of myocardial infarction, stroke, and coronary revascularization.

Primary Hyperlipidemia (including Heterozygous Familial Hypercholesterolemia): Repatha® is indicated as an adjunct to diet, alone or in combination with other lipid-lowering therapies (e.g., statins, ezetimibe), for the treatment of adults with primary hyperlipidemia to reduce low-density lipoprotein cholesterol (LDL-C).

Homozygous Familial Hypercholesterolemia: Repatha® is indicated as an adjunct to diet and other LDL-lowering therapies (e.g., statins, ezetimibe, LDL apheresis) for the treatment of patients with homozygous familial hypercholesterolemia (HoFH) who require additional lowering of LDL-C.

The safety and effectiveness of Repatha® have not been established in pediatric patients with HoFH who are younger than 13 years old or in pediatric patients with primary hyperlipidemia or HeFH.

Contraindication: Repatha® is contraindicated in patients with a history of a serious hypersensitivity reaction to Repatha®. Serious hypersensitivity reactions including angioedema have occurred in patients treated with Repatha®.

Allergic Reactions: Hypersensitivity reactions (e.g. angioedema, rash, urticaria) have been reported in patients treated with Repatha®, including some that led to discontinuation of therapy. If signs or symptoms of serious allergic reactions occur, discontinue treatment with Repatha®, treat according to the standard of care, and monitor until signs and symptoms resolve.

Adverse Reactions in Primary Hyperlipidemia (including HeFH): The most common adverse reactions (>5% of patients treated with Repatha® and occurring more frequently than placebo) were: nasopharyngitis, upper respiratory tract infection, influenza, back pain, and injection site reactions.

From a pool of the 52-week trial and seven 12-week trials: Local injection site reactions occurred in 3.2% and 3.0% of Repatha®-treated and placebo-treated patients, respectively. The most common injection site reactions were erythema, pain, and bruising.

Allergic reactions occurred in 5.1% and 4.7% of Repatha®-treated and placebo-treated patients, respectively. The most common allergic reactions were rash (1.0% versus 0.5% for Repatha® and placebo, respectively), eczema (0.4% versus 0.2%), erythema (0.4% versus 0.2%), and urticaria (0.4% versus 0.1%).

Adverse Reactions in the Cardiovascular Outcomes Trial: The most common adverse reactions (>5% of patients treated with Repatha® and occurring more frequently than placebo) were: diabetes mellitus (8.8% Repatha®, 8.2% placebo), nasopharyngitis (7.8% Repatha®, 7.4% placebo), and upper respiratory tract infection (5.1% Repatha®, 4.8% placebo).

Among the 16,676 patients without diabetes mellitus at baseline, the incidence of new-onset diabetes mellitus during the trial was 8.1% in patients assigned to Repatha® compared with 7.7% in those assigned to placebo.

Adverse Reactions in Homozygous Familial Hypercholesterolemia (HoFH): The adverse reactions that occurred in at least two patients treated with Repatha® and more frequently than placebo were: upper respiratory tract infection, influenza, gastroenteritis, and nasopharyngitis.

Immunogenicity: Repatha® is a human monoclonal antibody. As with all therapeutic proteins, there is potential for immunogenicity with Repatha®.

Please click [here](#) to see full Prescribing Information.

**RepathaReady® Program
Privacy Notice and Authorization**



In accordance with my signature below, I understand and consent to Amgen contacting me using the contact information provided in this form to enroll me in, operate, and administer Amgen patient support services and/or RepathaReady® program as described in the Patient Privacy Authorization other than promotional communications by telephone or SMS/text (to which I can separately opt-in below). I understand that the operation and administration of certain of these services and/or programs may require that Amgen contact me by telephone or SMS/text.

My preferred method(s) of contact:

Email Phone Mail SMS/text (standard text message charges may apply from your wireless provider)

In addition to the above consent, I understand that by checking this box and signing below, I consent to Amgen calling and texting me at the phone number(s) I have provided with promotional communications relating to Amgen products and services and/or my condition or treatment. Amgen may use automatic dialing machines or artificial or prerecorded messages to contact me and may leave a voicemail or text message (standard text messaging rates may apply). I understand that I am not required to provide this consent as a condition of purchasing any goods or services.

Amgen may contact me using the contact information provided in this form for participation in market research activities associated with Amgen's products, services and/or my condition or treatment. I agree / I disagree

My signature below certifies that I am at least 18 years old and that I have **read, understood, and agreed** to the Privacy Notice and Patient Authorization to release my personal health information as described in full detail on the next page.

Patient Name: _____

Name of Legal Guardian (if needed): _____

Patient Signature (or Legal Guardian): _____ Date: _____

Amgen's Privacy Pledge to Patients

Amgen respects patients and customers and takes the protection of their privacy very seriously. Amgen pledges the following:

- ✓ Amgen does not and will not sell or rent your information to marketing companies or mailing list brokers.
- ✓ Amgen is careful to only collect and/or use personal identifiable information for the purposes stated in this Authorization and, as necessary, to provide the services and/or programs the patient or customer chooses to enroll into.
- ✓ Amgen practices are consistent with federal and state privacy laws, including HIPAA.
- ✓ Amgen program enrollment is voluntary and always provides patients with an easy option to cancel participation.

Uses and Disclosure of Personal Information

I authorize Amgen and its contractors and business partners ("Amgen") to use and/or disclose my personal information, *including my personal health information, only for the following purposes:*

- To operate, administer, enroll me in, and/or continue my participation in Amgen's RepathaReady® program or any other Amgen-affiliated patient support services and activities related to my condition or treatment (e.g., co-pay card programs, reimbursement assistance programs, drug coverage verification, nurse educator services, adherence program, and disease management support);
- To contact, with my permission, my doctor and the rest of my healthcare team and share with them my health information that may be useful for my care;

RepathaReady® Program Privacy Notice and Authorization



Continued from previous page

- **To provide me with informational and promotional materials relating to Amgen products and services, and/or my condition or treatment;** and/or
- To improve, develop, and evaluate products, services, materials, and programs related to my condition or treatment.

In order for Amgen to provide me with the services and/or programs described above, Amgen needs to collect and use my personal information, including *my personal health information*. I understand that *my personal health information* may include any information, in electronic or physical form, in the possession of or derived from a healthcare provider, healthcare plan, pharmacy, pharmaceutical company, laboratory, and/or their contractor ("Healthcare Provider"). This may include select information from or about my medical history and general health, my healthcare plan benefits, payment limits or restrictions covered by my healthcare plan policy, and/or my adherence to my treatment.

I authorize my Healthcare Providers to disclose *my personal health information* to Amgen, and between themselves, as necessary, but only for the purposes stated above in this Authorization. I understand that certain of my Healthcare Providers (such as pharmacies and specialty pharmacies) may receive remuneration from Amgen in exchange for disclosing *my personal health information* and/or for using my information to contact me with communications about Amgen products that have been prescribed to me (e.g., adherence programs) and other patient support services.

Expiration, Right to Obtain a Copy, and Right to Cancel

I understand that by signing this form, I authorize my Healthcare Providers or others who might hold my health information to only release it to Amgen employees, as well as to its contractors and business partners, who are performing the services set forth in this Authorization. I also understand I am authorizing my personal information, including *my personal health information*, to be used for the purposes described above. I understand and agree that by signing below, I am authorizing those who rely on this Authorization to release my personal health information for the earlier of five (5) years or until my participation in the program ends through my cancellation, unless a shorter time period is required by state law.

I understand that I can obtain a copy of this Authorization or cancel this Authorization at any time by calling Amgen at 1-844-REPATHA or by writing to PO Box 220326; Charlotte, NC 28222. If I cancel my consent, I will no longer qualify for the services described. I also understand that if a Healthcare Provider is disclosing my personal health information to Amgen on an authorized ongoing basis, my cancellation with Amgen will be effective with respect to any such Healthcare Providers as soon as they receive notice of my cancellation.

No Effect on Treatment

I understand I do not have to sign this Authorization and that my enrollment in any of the services and/or programs described above is entirely voluntary. I understand that Amgen, as well as Healthcare Providers, cannot require me, as a condition of having access to medications, prescription drugs, treatment or other care, to sign this Authorization. Federal Law (including HIPAA) requires a signed authorization in order for Amgen to collect this information from my Healthcare Providers. I understand I cannot participate in the listed services and/or programs without signing this Authorization or an equivalent authorization with my Healthcare Providers.

Information Received From Healthcare Providers

I understand that once my personal health information has been disclosed to Amgen, federal privacy laws may no longer apply and protect it from further disclosure. Amgen agrees, however, to protect my personal health information by only using and disclosing it as stated in the Authorization or as otherwise allowed or required by law. I understand that Amgen does not and will not sell or rent my information to marketing companies or mailing list brokers.

