



- Partner with and remain dedicated to the patient throughout their treatment journey
- Contact the patient or caregiver to review insurance coverage and support programs

For Patients

PATIENT CONSENT TO COLLECT, USE AND SHARE PERSONAL INFORMATION (PI) AND SIGNATURE

(Sections 1–4 to be read and completed by Patient or Patient’s Authorized Representative)

The purpose of this form is to permit HoFH patients who have been prescribed EVKEEZA® (evinacumab for injection) to receive information and support (“Patient Support”) from UltraCare, its affiliates, representatives, agents, and contractors. UltraCare® provides Patient Support to eligible patients who have been prescribed EVKEEZA. This includes: (1) providing reimbursement and financial support to eligible patients (such as investigating your insurance coverage and reviewing eligibility for financial assistance); (2) working with you and your provider to fill your prescription; (3) providing you with disease- and medication-related educational resources and communications; and (4) coordinating EVKEEZA infusions.

The UltraCare® Program (“Program”) is sponsored by Ultragenyx Pharmaceutical, Inc. (“Ultragenyx”) and administered by Innomar on behalf of Ultragenyx.

Please read this form carefully and ask any questions that you may have before signing.

1. PATIENT INFORMATION (Be sure to choose your preferred contact method)

| | |
|---|---|
| First, Middle, Last Name _____ | Street Address _____ |
| Gender <input type="checkbox"/> Female <input type="checkbox"/> Male <input type="checkbox"/> Other DOB (DD/MM/YYYY) _____ | City _____ |
| Health Card Number _____ | Province _____ Postal Code _____ |
| Home Phone (_____) _____ Work Phone (_____) _____ | Email _____ |
| Mobile Phone (_____) _____ Best Time to Contact _____ | Caregiver Name (First and Last) _____ |
| Preferred Method of Contact: <input type="checkbox"/> Home <input type="checkbox"/> Work <input type="checkbox"/> Mobile <input type="checkbox"/> Email | Relationship to Patient _____ Caregiver Phone (_____) _____ |
| Preferred Language: <input type="checkbox"/> English <input type="checkbox"/> French <input type="checkbox"/> Other _____ | |

2. AUTHORIZATION TO SHARE PROTECTED HEALTH INFORMATION

By signing below, I authorize my healthcare providers, including my physicians and pharmacies (“My Providers”) and my health insurance plan (“My Plan”) to share my medical information (such as information about my diagnosis, prescriptions, and treatment) and my insurance information (“My Information”) with UltraCare® so that UltraCare® can provide Patient Support. I authorize My Providers to use My Information to provide me with certain offerings related to my medication and treatment. I understand that once My Information has been disclosed, federal privacy laws may no longer protect the information. However, I understand that UltraCare® agrees to protect My Information by using and disclosing it only for purposes described in this Authorization or as required by law or regulations. I understand that Innomar deploys security practices that include the development of internal controls that restrict both logical and physical access to its systems to appropriate authorized personnel. As a component of Data Security, Innomar invests in data encryption per industry regulations and standards. Data is encrypted across all company assets, including servers and workstations. I understand that these efforts will be made to keep my PI confidential.

I understand that I may refuse to sign this Authorization, and that my treatment, insurance enrolment, and eligibility for insurance benefits are not conditioned upon signing this Authorization. I also understand, however, that refusing to sign this Authorization means that I may not participate in UltraCare® and may not be able to take advantage of other offerings by UltraCare®. I may cancel or revoke this Authorization at any time by letting UltraCare® know. I understand that if I revoke this Authorization, My Providers and UltraCare® will stop using and sharing My Information under this Authorization, and additional information will not be collected. My revocation will not affect uses and disclosures of My Information prior to my revocation. I understand that unless my consent is withdrawn, my consent is valid for the longer of either ten (10) years from the date signed below or as long as I receive services from the Program and for a reasonable time thereafter. I understand that I may receive a copy of this Authorization.

| | | | |
|---|---|-------------------------|-------|
| _____ | X | _____ | _____ |
| Print Patient or Authorized Patient Representative Name | Signature of Patient or Authorized Patient Representative | Relationship to Patient | Date |

3. AUTHORIZATION FOR ULTRACARE® AND COMMUNICATIONS

By signing below, I confirm I would like to enrol in the UltraCare® program and authorize UltraCare® to provide me with Patient Support. I understand that UltraCare® is an optional program. I agree that UltraCare® may use My Information and share it with My Providers or My Plan in connection with providing the Patient Support, administering the UltraCare® program, or as otherwise required by UltraCare® to meet its legal obligations. For example, UltraCare® may communicate with me (such as by mail, phone, email, and/or text message) or my caregiver, use My Information to tailor the UltraCare®-related communications to my needs, and share information with My Providers about dispensing my EVKEEZA medicine to me. I understand that UltraCare® may de-identify My Information, combine it with information about other patients, and use the resulting information for UltraCare® reporting purposes.

| | | | |
|---|---|-------------------------|-------|
| _____ | X | _____ | _____ |
| Print Patient or Authorized Patient Representative Name | Signature of Patient or Authorized Patient Representative | Relationship to Patient | Date |

4. OPT IN TO RECEIVE MARKETING COMMUNICATIONS (NOT REQUIRED FOR ULTRACARE® ENROLMENT)

By checking this box, I authorize UltraCare®, and companies working with UltraCare®, to contact me by mail, email, fax, and/or telephone regarding marketing and promotional communications, customer surveys, or for market research surveys. I understand that I am not required to provide this consent to receive marketing communications to receive EVKEEZA or UltraCare® services.

| | | | |
|--|--|--------------------------------|-------------|
| | X | | |
| Print Patient or Authorized Patient Representative Name | Signature of Patient or Authorized Patient Representative | Relationship to Patient | Date |

IMPORTANT: If healthcare provider is unable to obtain written consent from patient, please document when patient verbal consent was obtained. This will allow the program to continue with processing this enrolment. Written consent will be obtained by the program. Verbal consent should be obtained by a healthcare provider.

Patient consented verbally: Date (DD/MM/YYYY) _____

Patient consent obtained by: Name (First, Last) _____ Title: MD RN Other (specify) _____

Signature _____

By providing my email address, I agree to receive, electronically, communications from Innomar acting on behalf of Ultragenyx Pharmaceutical, Inc. containing information and updates relating to enrolment in the UltraCare® Program.

I understand that I may withdraw any of my consents to such communications at any time by providing notice to Innomar Strategies, Inc., c/o UltraCare® Program, 2600 Alfred Nobel Blvd, Ville Saint-Laurent, QC H4S 0A9, or via email at Ultracare@innomar-strategies.com

You can report any suspected side effects associated with the use of health products to Health Canada at 1-866-234-2345 or <http://www.hc-sc.gc.ca/dhp-mps/medeff/report-declaration/index-eng.php>. You may also report side effects to Ultragenyx at 1-833-388-5872 (U-LTRA).

FOR HEALTHCARE PROVIDERS

(Sections 5-9 to be read and completed by Healthcare Provider)

Patient Name (First, Last) _____

DOB (DD/MM/YYYY) _____

5. PRESCRIBER INFORMATION

| | |
|------------------------------------|----------------------------------|
| First name _____ | Street address _____ |
| Last name _____ | City _____ |
| Office email _____ | Province _____ Postal code _____ |
| Office contact name/title _____ | Office phone (_____) _____ |
| Office contact phone (_____) _____ | Fax (_____) _____ |
| Licence # _____ | Prescriber email _____ |

6. CONFIRMED DIAGNOSIS OF HOMOZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA (HOFH) BY:

Clinically diagnosed Genetic confirmation of bi-allelic pathogenic/likely pathogenic variants on different chromosomes at the LDLR, APOB, PCSK9, or LDLRAP1 genes Confirmed diagnosis of OTHER _____

7. PATIENT HISTORY

Patient Status and History

Current LDL-C value (pre-apheresis if applicable)

_____ mmol/L Date (DD/MM/YYYY) ____/____/____

Untreated LDL-C value (prior to treatment initiation)

_____ mmol/L Date (DD/MM/YYYY) ____/____/____

Cutaneous or tendinous xanthoma Age of xanthoma onset _____

Family History

Evidence of HeFH in both parents

Lipid-Lowering Treatments

| | Treatment name | Dose | Current | Previous | Duration of treatment |
|--------------------------|------------------------|-------|--------------------------|--------------------------|-----------------------|
| <input type="checkbox"/> | Statin | _____ | <input type="checkbox"/> | <input type="checkbox"/> | _____ |
| <input type="checkbox"/> | Ezetrol® (ezetimibe) | _____ | <input type="checkbox"/> | <input type="checkbox"/> | _____ |
| <input type="checkbox"/> | PCSK9i | _____ | <input type="checkbox"/> | <input type="checkbox"/> | _____ |
| <input type="checkbox"/> | Juxtapid® (lomitapide) | _____ | <input type="checkbox"/> | <input type="checkbox"/> | _____ |
| <input type="checkbox"/> | Other | _____ | <input type="checkbox"/> | <input type="checkbox"/> | _____ |

Lipoprotein apheresis or Plasmapheresis weekly bi-weekly monthly other _____

8. INFUSION SETTING AND ADMINISTRATION

Preferred Treatment Setting

Out-patient clinic* Apheresis unit* * Provide contact name and phone number.

Innomar clinic At home

Contact Name _____ Phone Number _____

9. EVKEEZA (EVINACUMAB FOR INJECTION) FOR INFUSION USE – PRESCRIPTION INFORMATION

The recommended dose for EVKEEZA is 15 mg/kg administered by intravenous (IV) infusions over 60 minutes every 4 weeks. The rate of infusion may be slowed, interrupted, or discontinued if the patient develops any signs of adverse reactions, including infusion-associated symptoms. EVKEEZA can be administered without regard to lipoprotein apheresis.

| | |
|---|---|
| <p>REQUIRED Patient's Full Name _____</p> <p>Patient weight in kg _____</p> <p>Date _____</p> <p>Dose: 15 mg/kg IV every 4 weeks according to the weight of the day</p> <p>Special instructions/Indication: Administer by intravenous infusion over 60 minutes</p> | <p>Infusion fluid type (please select one):</p> <p><input type="checkbox"/> 0.9% Sodium Chloride Injection</p> <p>or</p> <p><input type="checkbox"/> 5% Dextrose Injection</p> <p>Refills _____ Days' supply: 4 weeks</p> |
|---|---|

If patient has already started treatment, EVKEEZA supply needed for scheduled treatment on (DD/MM/YYYY) ____/____/____

How Supplied: EVKEEZA (evinacumab for injection) is supplied by 150 mg/mL concentrate for solution for infusion (DIN: 02541769).

Please see full Product Monograph at <https://www.canada.ca/en/health-canada/services/drugs-health-products/drug-products/drug-product-database.html> for complete dosage and administration information.

Prescriber Signature _____ Date _____

Special Instructions _____

Special Precautions (e.g., allergies) _____

The prescriber assumes responsibility for monitoring lab values. The prescriber assumes responsibility for notifying UltraCare® of any dosage changes or suspension of therapy.