

PATIENT ENROLLMENT AND PRESCRIPTION FORM



Fax: 877-914-0660 | Phone: 844-789-8744 Hours of Operations: Monday - Friday, 8 AM to 8 PM ET

Complete this form for each patient, fields in red with asterisks are required.* PATIENT INFORMATION Patient First Name*: _____ Last Name*: _____ DOB*: __/__/ Gender:

Male Female _____ City*: _____ Street Address*: _____ _____ State*: _____ ZIP Code*: _____ Alternative #: Primary Phone #* (mobile preferred): _____ Language: English Spanish Other Alternate Contact / Care Partner Full Name: __ Relationship to Patient: ___ _____ Email: _____ Phone #: PRESCRIBER INFORMATION Prescribing Provider Name*: _____ _____ Specialty: _____ Lipid Specialty Office Contact Name: _____ Practice Name: _____ State*: _____ ZIP Code*: ____ Practice Address*: _____ City*: ____ Office Phone #*: Office Fax #*: ______ Email: ____ Prescribing Provider NPI #*: 3 DIAGNOSIS AND CLINICAL INFORMATION Attach copy of triglyceride tests, past and current medications, drug allergies, and acute pancreatitis history Indication*: Familial Chylomicronemia Syndrome (FCS) Other ICD-10-CM Diagnosis Code: ☐ E78.3 Hyperchylomicronemia ☐ Other TRYNGOLZA™ PRESCRIPTION INFORMATION Prescriber Instructions: Comply with State-specific requirements such as e-prescribing, state specific prescription form, fax language, etc. Either (1) fill out the information below and provide signature, or (2) send the prescription electronically to PANTHERX Specialty Pharmacy Rx: TRYNGOLZA™ (olezarsen) injection, 80 mg/0.8 mL single-dose autoinjector 10-digit NDC: 71860-101-01 Dosing: ☐ Administer 80 mg subcutaneously once monthly OR ☐ Other dosing instructions: Quantity = QS for 30 days supply OR Other quantity: ______ Refills:___ Dispense as written Quick Start: If eligible and when all information required for prior authorization is received, patient will be enrolled in Quick Start program that will provide free drug during the insurance approval process. The Quick Start program is available to all insured patients who are US residents with a diagnosis of familial chylomicronemia syndrome (FCS). Eligibility is subject to the terms and conditions of the program. Ionis Pharmaceuticals® reserves the right to rescind, revoke, or amend the program at any time without notice. I request for my patient **NOT** be provided Quick Start By signing this form, I am indicating a prescribing decision has been made. In addition, I am certifying treatment with TRYNGOLZA™ indicated above is medically necessary for this patient, and I have received authorization to release the medical and/or other patient information relating to this therapy to Ionis Every Step™ and its affiliates, agents, and representatives to use and disclose as necessary for prior authorization processing and fulfillment of the prescription. I certify that, to the best of my knowledge, the patient and physician information in this form is complete, accurate, and consistent with applicable privacy regulations. For Quick Start: I understand that this medication is being provided free to the named patient by Ionis and agree that neither I nor the patient will bill an insurer or any government healthcare program for the cost of this medication. The program may not be combined with another offer and is not eligible to patients without insurance or whose insurer has made a final coverage determination. MM/DD/YYYY Prescriber Signature (Physician attests this is his/her legal signature. NO STAMPS)

Please see Indication & Important Safety Information on page 3 and full Prescribing Information for TRYNGOLZA, also available at TRYNGOLZAhcp.com

Attention: NY providers, please submit electronic prescriptions.



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NSURANCE INFO	RMATION		
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EVERY STEP

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INDICATION & IMPORTANT SAFETY INFORMATION FOR TRYNGOLZA

INDICATION

TRYNGOLZA (olezarsen) is indicated as an adjunct to diet to reduce triglycerides in adults with familial chylomicronemia syndrome (FCS).

IMPORTANT SAFETY INFORMATION

CONTRAINDICATIONS

TRYNGOLZA is contraindicated in patients with a history of serious hypersensitivity to TRYNGOLZA or any of the excipients in TRYNGOLZA. Hypersensitivity reactions requiring medical treatment have occurred.

WARNINGS AND PRECAUTIONS

Hypersensitivity Reactions

Hypersensitivity reactions (including symptoms of bronchospasm, diffuse erythema, facial swelling, urticaria, chills, and myalgias) have been reported in patients treated with TRYNGOLZA. Advise patients on the signs and symptoms of hypersensitivity reactions and instruct patients to promptly seek medical attention and discontinue use of TRYNGOLZA if hypersensitivity reactions occur.

ADVERSE REACTIONS

Most common adverse reactions (incidence >5% of TRYNGOLZA-treated patients and >3% higher frequency than placebo) were injection site reactions, decreased platelet count, and arthralgia.

Please see full Prescribing Information for TRYNGOLZA

