Welcome to the **Navigator Program**



Start Form

A no-cost program created to provide access support to your patients throughout their entire treatment journey with CUVRIOR[®] (trientine tetrahydrochloride)

From start to finish, you can rely on your dedicated Care Coordinator to be there and help your patients

- Get started on therapy while you and the pharmacy work with their insurance plan to obtain ongoing coverage
- Understand their insurance coverage for CUVRIOR •
- Understand the processes required by their insurance to access CUVRIOR (such as prior • authorizations)
- Receive information related to co-pay and/or financial assistance programs that may be available

Get started

There are 2 ways to get started in the pursuit of CUVRIOR:

- Submit a completed Start Form to enroll your patient in the Navigator Program. With this form, 1 the patient will be automatically evaluated for program offerings that they may be eligible for.
- Submit an eRX or provide a verbal Rx to PANTHERx. This option starts the Benefits Investigation 2 process, but does not specifically enroll the patient in the program. If the patient wants to take advantage of offerings, additional outreach to the physician may be required in order to obtain the appropriate attestations.

Ready to enroll?



👔 🕤 Have questions about the Navigator Program? Patients may contact a Care Coordinator today to discuss how the Navigator Program can support you throughout your treatment journey: 1-877-995-ORPH (6774)



Navigator Program Start Form

Submit Completed Start Form to Navigator Program

Phone #: 1-877-995-ORPH (6774)

Fax #: 1-866-716-ORPH (6774)

Navigator Program[™]

Patient Support by Orphalan

Step 1 Patient Information	on						
First Name:	Middle Initial:		Last Name:				
Date of Birth:					Sex: O Male O Female		
Primary Address:							
City:		State: ZIP:					
Preferred Phone #:	(Home or mobile):	me or mobile):		OK to Leave Message: O Yes O No			
Alternate Phone #:		Primary Email:					
Caregiver Name:		Caregiver Phone #:					
Relationship to Patient:							
Step 2 Insurance Inform	ation *Please include	front & back	copies of all	insurance ca	ards (<i>medical & pharmacy</i>)*		
PHARMACY BENEFIT INFORMATION:							
Prescription Insurance Carrier:		PBM Ph	one #:				
Member ID #:							
BIN #: Policy Holder Name:							
Secondary Prescription Carrier:		_ PBM Phone #:					
Member ID #:							
	\ #:						
	Holder Name:						
MEDICAL INSURANCE INFORMATION:		locurao	co Dhono #:				
-	Medical Insurance:						
		_ Group I	u #				
Insurance Type:							
O Commercial O Medicare O Medic							
Policy Holder Name:		_ Relatior	nship to Patie	nt:			
Secondary Medical Insurance:		_ Insuran	ce Phone #: _				
Member ID #:		_ Group ID #:					
Insurance Type:							
O Commercial O Medicare O Medic	aid 🔿 TRICARE/DoD)		

Step 3 Diagnosis and Clinical Information							
ICD/Diagnostic Code(s): O Wilson Disease (E83.01) O Other Diagnosis:							
Current Therapy for Wilson Disease:							
	Dose:		Duration:				
Medication:	Dose:		Duration:				
Previous Therapies (Check all that apply):			Medication Allergies	s:			
○ Cuprimine [®] ○ D-Penamine [®]	O Depen Titratabs®	O Penicillamine (generic)	🔘 Known Drug Aller	gies:			
○ Syprine [®] ○ Trientine (<i>generic</i>)	◯ Galzin®	⊖ Wilzin®					
O OTC Zinc O CUVRIOR		O Other:	No Known Drug A	Allergies			
	-	-	-				
Step 4 Prescription Info	rmation *Each pres	cription must be completed in	its entirety for the pres	cription to be valid*			
First Name:	Middle Initial: Last Name:						
Date of Birth:	Sex: O Male O Female						
Quick Start Program: an optional program	available to eligible patie	nts at no cost (See Terms of Par	ticination on page 5)				
Quick Start Program: an optional program	available to eligible patie		ucipation on page 5)				
Prescriber Attestations (<i>Required for participation in the Quick Start Program</i>): I confirm that the patient has been diagnosed with Wilson Disease, is age 18 or older, is not treatment naïve (patient has been de-coppered), and has not previously been treated with CUVRIOR.							
I understand my patient's eligibility and receipt of free product is not contingent on any purchase obligations from Orphalan, its agents or service providers. I also understand any products distributed under the Navigator Program are provided free of charge, and may not be submitted for reimbursement to any payor, including a federal healthcare program, and may not be sold, traded, distributed for sale or returned for credit; nor may I bill for administration of such product. I agree to assist in efforts to secure coverage for CUVRIOR for my patient.							
Quick Start Prescription: CUVRIOR® (trientine tetrahydrochloride) Dispense Quantity:	-	-day supply Refill(s): 1 refill	Directions for Use:				
Ongoing Prescription:							
CUVRIOR [®] (trientine tetrahydrochloride)	300 mg Tablets						
Dispense Quantity: Days' Supply:			Refills:				
Directions for Use:							
Step 5 Prescriber Inform	nation						
	Prescriber Last Name:						
Prescriber Specialty:							
Street Address:							
	Prescriber Fax:						
	Office Contact Title:						
Office Contact Phone: Office Contact Email:							
Prescriber Authorization:							
I certify that I am the prescriber mentioned necessity information included on this Start F prescribed CUVRIOR therapy for an FDA-appr obtained the patient's written authorization to administer the Navigator Program, to Orphala PANTHERx Rare, as the dedicated specialty ph to the patient's insurance plan if permitted by I am licensed to prescribe the product listed of practicing in a state with official prescription r	orm and attestations (<i>as a</i> , oved indication and I will b o provide the information an, its agents and service p narmacy agent and on beh y the policies of that plan a on this form and the prescu	<i>pplicable</i>) are true, accurate, and be supervising the patient's trea in this Start Form, and such oth providers in accordance with all half of my patient, to (1) forward nd (2) coordinate medication de ription complies with my state-s	d complete. I further cer tment accordingly. I cert er information as may b applicable federal and s the above statement of elivery with the patient. F specific prescribing requi	tify that I have ify that I have e necessary to tate laws. I authorize medical necessity Finally, I certify that			
Prescriber Signature:	•	Dispense as Written 🔘 Substi	tution Allowed Date: _				
(Signature stamps o	are not acceptable)						
Please see Quick Start Program Terms of Part Please see Important Safety Information on p © 2024 Orphalan. All rights reserved. US-ORP	age 6 and full Prescribing	Information on www.cuvrior.c	com.	Page 3 of 6			

Step 6

AUTHORIZATION FOR USE AND DISCLOSURE OF PERSONAL INFORMATION

I would like to enroll in the Navigator Program. By signing this Enrollment Form ("Authorization") I authorize Orphalan, Inc., and its affiliated companies, agents and service providers (collectively, "Orphalan") to provide me with support under the Navigator Program.

I authorize Orphalan, my health care providers, and their staff ("HCPs"), my health insurer(s), patient assistance organizations, and my pharmacy providers ("Authorized Parties") to use, process and share: (1) my personal health information (e.g., information about my diagnosis, treatment and medical condition), (2) information that identifies me (e.g., my name, address, phone number, date of birth), and (3) my insurance information (collectively my "Personal Information").

I understand this sharing of my Personal Information is necessary to enable the Authorized Parties to enroll me in the Navigator Program, provide Navigator Program services to me, operate the Navigator Program, conduct other business activities, and meet legal requirements. For example, Orphalan must use my Personal Information to communicate with me, investigate insurance matters, determine my eligibility for patient support services, and coordinate with my HCP about my enrollment. I understand that Orphalan may use "de-identified" data from the Navigator Program, meaning it may remove elements of my Personal Information that identify me, combine my data with other patients' information and use this "de-identified" data for business purposes such as analysis and reporting. I understand that once my Personal Information is shared, federal privacy laws may no longer protect it, and may not prevent re-disclosure by Orphalan or the Navigator Program. However, I understand Orphalan and the Navigator Program have agreed to use, process and disclose my Personal Information only for the purposes described in this Authorization. For more information about how Orphalan collects, uses, and protects my Personal Information, I can visit www.cuvrior.com to review the Privacy Policy.

I understand the Navigator Program is optional. If I choose not to sign this Authorization, I am still able to receive the medication that has been prescribed to me by my HCP. But not signing this Authorization will mean I cannot participate in the Navigator Program.

This Authorization expires five (5) years from the date of my signature below unless a shorter period is required by state law or I cancel my participation in the Navigator Program.

California Residents: I have the right to access my Personal Information, update my Personal Information if it is incorrect, or to request that Orphalan delete or limit the use of my Personal Information. I understand that deletion may not be possible or required under certain circumstances. To exercise the use of this right, I must send a written notice by mail to the address provided below.

I understand that I may revoke this authorization by sending a written notice to the Navigator Program at **Navigator Program**, 24 Summit Park Dr., Pittsburgh, PA 15275 or faxing a written request to 1-866-716-6774. I understand that if I revoke this Authorization, my revocation will not invalidate any uses or disclosures of my Personal Information made in reliance on this Authorization prior to the Navigator Program's receipt of this notice.

By signing below, I acknowledge that the Navigator Program provided by Orphalan will make payments to third-party providers for processing my Personal Information. I also understand that each patient support offering under the Navigator Program is subject to its own terms and conditions, and these terms and conditions may change over time. If I am eligible to receive one or more of the Navigator Program's patient support offering(s), I will be required to abide by all applicable terms and conditions; and I agree to do so. I acknowledge that my eligibility for each patient support offering under the Navigator Program depends on my insurance situation, and if my insurance information changes at any time, I will notify the Navigator Program as soon as possible.

By signing below, I certify that I have read and agree to the above.						
Patient's Name (<i>Printed</i>):	Date:					
Patient, or Personal Representative Signature:						
Personal Representative's Description of Authority (<i>If applicable</i>):						
Call and Text Opt-in (<i>Optional</i>): By checking this box, I further an number(s) I provide.* I understand I may revoke my consent to re text from the Navigator Program or by contacting the Navigator F	eceive automated calls or text messages by replying STOP to any					
Marketing Opt-in (<i>Optional</i>): I authorize Orphalan to contact me by mail, email, fax, and/or telephone regarding other potential topics of interest to me, customer surveys, or occasionally for market research purposes. I understand that I am not required to provide this consent as a condition of receiving any Orphalan medicine or patient support services.						
Patient Phone #:	_ Patient Email Address:					
*I understand texts I receive may be subject to fees imposed by my tele	communications provider, and I will be responsible for paying these fees.					

Step 7 QUICK START TERMS OF PARTICIPATION

The Quick Start Program (Quick Start) is available for insured patients who are U.S. residents, age 18 and older, new to CUVRIOR[®], and experiencing a coverage delay that has lasted at least five days. Additional eligibility criteria may apply. Quick Start provides a 30-day supply of CUVRIOR[®] to eligible patients while they work with their health care provider (HCP) and insurer to obtain coverage. A one-month refill is available if coverage delays persist. Patients and HCPs have a responsibility to pursue coverage diligently. Patients may be asked to reverify their insurance coverage status during the course of their participation in Quick Start.

Quick Start is not insurance, nor is participation a guarantee that insurance coverage will be obtained successfully. Participation in Quick Start does not require the patient or prescriber to make any future purchases from Orphalan or PANTHERx. Free product received under Quick Start may not be submitted to any third-party payor for reimbursement, and may not be sold, traded or distributed to anyone other than the patient for whom it was intended. For Medicare Patients: Quick Start product is administered outside the Medicare benefit and should not be counted toward the patient's true out-of-pocket ("TrOOP") cost for any Part D enrollee.

Participation in Quick Start concludes at the earlier of successful coverage or exhaustion of permitted refills. Patients may not re-enroll. Orphalan may change or terminate Quick Start at any time without notice.

CUVRIOR[®] (trientine tetrahydrochloride) Tablets

INDICATION

CUVRIOR is a prescription medicine used to treat adults with stable Wilson disease who are de-coppered and able to take penicillamine. Wilson disease is a condition where the body stores too much copper. CUVRIOR is a chelating medicine which binds copper in the blood stream and eliminates it in the urine to help reduce excess copper in the organs.

IMPORTANT SAFETY INFORMATION

- **Do not use** CUVRIOR if you are allergic to trientine or to any of the ingredients in CUVRIOR.
- Your Wilson Disease symptoms could get worse when you start treatment. This could happen because too much copper is removed from the body in a short period of time. Your doctor may need to reduce your dose or stop CUVRIOR treatment.
- **Copper deficiency** may develop following treatment with CUVRIOR. Your doctor will do tests to monitor your urine and blood for copper.
- **Iron deficiency** may develop while taking CUVRIOR. If this happens, your doctor may tell you to take iron supplements for a limited time.
- Allergic reactions, such as a rash can occur. If a rash or other allergic reaction occurs, contact your doctor immediately or get emergency help.
- The most common side effects are pain in the abdomen, change in bowel habits, rash, hair loss (alopecia), and mood swings.

Take CUVRIOR at least 1 hour apart from any other oral medicine. Tell your doctor about all the medicines you're taking, including over-the-counter medicines, vitamins, and herbal supplements. Avoid taking supplements when taking CUVRIOR. Taking CUVRIOR with mineral supplements (e.g., iron, zinc, calcium, magnesium) can reduce the effectiveness of CUVRIOR. If iron supplementation is necessary, take CUVRIOR at least 2 hours before or 2 hours after taking iron; for other mineral supplements (e.g., zinc, calcium, magnesium), take CUVRIOR at least 1 hour before or 2 hours after. Do not start or stop any medicines while taking CUVRIOR without talking to your doctor.

These are not all the possible side effects of CUVRIOR. You are encouraged to report negative side effects of prescription drugs. Contact Orphalan at 1-800-961-8320 or FDA at 1-800-FDA-1088 or **www.fda.gov/medwatch**.

Please see full Prescribing Information at www.cuvrior.com.