

PROCYSBI® (CYSTEAMINE BITARTRATE) DELAYED-RELEASE CAPSULES AND DELAYED-RELEASE ORAL GRANULES PATIENT ENROLLMENT FORM INSTRUCTIONS

The Patient Enrollment Form is required to initiate treatment with Horizon Therapeutics' prescription medicine, PROCYSBI.

Instructions:

- 1. Fill out all patient information, including the most recent results of a white blood cell (WBC) cystine level test, recent history with CYSTAGON® (cysteamine bitartrate) capsules, and use of a gastrostomy tube (G-tube).
- **2.** Fill out all required prescriber information, including all contact information for the practice or facility.
- **3.** Complete and/or review all required insurance information for the patient and, if possible, attach copies of the patient's insurance cards for primary as well as supplementary insurance.
- **4.** Complete the prescription and clinical information in its entirety; all fields are required. Reference the included select PROCYSBI dosing instructions or the PROCYSBI Full Prescribing Information for complete dosing information.
- **5.** Review, sign, and date the prescriber certification at the bottom of the Patient Enrollment Form. In signing, you are indicating to dispense PROCYSBI as written. If a substitution is allowed, it should be noted.
- **6.** Check with your patient to ensure he or she has printed, signed, and dated the required Patient Authorization Form providing HIPAA authorization for Horizon By Your Side, a patient support program, and initiation of patient support.
- **7.** Fax pages 1 and 2 of this form, along with both sides of the patient's medical and prescription drug benefit cards, to the Horizon By Your Side team at 1-877-773-9411, or email them to PROHBYS@horizontherapeutics.com. Retain a copy of this form in the patient's records.

Please see IMPORTANT SAFETY INFORMATION on last page and <u>click here for the PROCYSBI Full Prescribing Information</u>.





PROCYSBI® (CYSTEAMINE BITARTRATE) DELAYED-RELEASE CAPSULES AND DELAYED-RELEASE ORAL GRANULES PATIENT ENROLLMENT FORM



Please fax completed form to 1-877-773-9411, or email it to PROHBYS@horizontherapeutics.com.

Phone: 1-855-888-4004 Fax: 1-877-773-9411 <u>PROCYSBI.com</u>

1. PATIENT INFORMATION						
First Name	MI		Last Name			
Address		City		State_	Zip	
Home Phone		Mobile Pho	ne			
Date of Birth		Gender 🔲	M 🔲 F Height		Weight	
Email		Preferred N	Nethod of Contact Hom	ne 🗌 Mob	ile 🗌 Email 🔲 Mail	
Currently taking CYSTAGON® (cyst	reamine bitartrate)?	No Last CYSTA	.GON daily dose (mg/day) _			
Currently on dialysis? ☐ Yes ☐ N	lo	Does the pa	atient have a G-tube (feedin	ig tube)? [□Yes □No	
White blood cell (WBC) test in the	last year? ☐Yes ☐No	(A bolus [st	raight] feeding tube 14 Fren	nch or large	er is recommended.)	
ALTERNATIVE CONTACT AND/OR	CAREGIVER					
First Name	Last Name		Hor	ne Phone .		
Mobile Phone	Email		Preferred Method	of Contact	Home Mobile Emai	
2. PRESCRIBER INFORMATION	ON		Preferred Method	of Contact	Email Phone	
Prescriber First Name	MI Las	st Name	Prescr	iber NPI# .		
Address						
Phone	Fax		Physician Specialty			
Office Contact Name		Email		_ Phone _		
3. INSURANCE INFORMATION	N — Please attach a copy of both	sides of the patier	ıt's insurance card(s).		■ No Insurance	
PRIMARY INSURANCE		SECON	DARY INSURANCE (if any)			
Insurance Carrier		Insurar	nce Carrier			
Customer Service Phone		Custor	ner Service Phone			
Subscriber Name		Subscr	iber Name			
Patient's Relationship to Subscribe	Patient	Patient's Relationship to Subscriber				
Subscriber Date of Birth Subsc			Subscriber Date of Birth			
Subscriber ID Number	riber ID Number Subscriber ID Number					
Policy/Employer/Group Number	Number Policy/Employer/Group Number					
Prescription Card? ☐ Yes If Yes, C	n Card? Yes If Yes, Carrier: Phone					
4. PRESCRIPTION AND CLINI	CAL INFORMATION					
Diagnosis (ICD-10-CM Code)	E72.04 Other					
Drug Name: PROCYSBI Capsule	es: 25 mg Quantity	v and/or □75 m	gQuantity		eg, Capsules: 600 mg q12h or 500 mg (6 x 75 mg capsules + 2 x 25 mg capsules) q12 Packets: 600 mg q12h or 525 mg (1 x 300 mg packets + 3 x 75 mg packets) q12h.	
•			•	i		
Directions: Days' Suppl	y Refills			D	lose Titration, see PROCYSBI Dosing	
Drug Name: PROCYSBI Granule	Packets: □75 mg C	uantity and/or	7300 mg Ou	Ir	Information for Healthcare Prescribers on page 3 for more information.	
_	_	-	_	-	Note: The prescriber is to comply with his/he state-specific prescription requirements such as e-prescribing, state-specific prescription form, fax language, etc.	
Directions: Days' Supply	Refills		,	a fo		
s the patient allergic to penicillamine	e, cysteamine, or any other medica	tion? If yes, please	e list:			
Prescriber Certification certify that the above therapy is medically necessary, that the int dist affiliates and their respective employees or agents (col on-medical treatment support for PROCYSBI, as prescribed, a tealth information with Horizon for purposes of the Program an ROCYSBI and assistance in initiating or continuing PROCYSBI may express or implied agreement or understanding that I wou nedical necessity; and (c) I will not seek reimbursement for an he completion and submission of coverage- or reimbursement state requirements: The prescriber is to comply with his/her state to the program of the process he services and support offered by Horizon By Your Side un attent to determine whether the patient is interested in signature.	lectively, "Horizon") will use this information to administ min educating about the insurance process. By my signar d (2) I have obtained the patient's authorization to releas as prescribed. I further understand and agree that (a) at ald recommend, prescribe, or use PROCYSBI or any other by medication or service provided by or through the Prognt-related documentation are the responsibility of the pae-e-specific prescription requirements such as e-prescribing in Horizon By Your Side has initiated; however, your nless your patient signs a Patient Authorization, cons	er the Horizon By Your Side pr ture, I also certify that (1) my le e such information as may be by medication or service provior I Horizon product or service, fram ram from any government pro tient and healthcare provider, state-specific prescription for batient must sign a Patient A	ogram (the "Program"), which provides a wide patient or his/her personal representative has p required for Accredo Health Group, Inc. (or anot ded through the Program as a result of this for for any other person; (b) my decision to prescril gram or third-party insurer. I understand that H Horizon makes no representation or guarantee m, fax language, etc. Noncompliance with state-s- uthorization to complete enrollment in Horiz	array of patient-from the party acting or m is for the name the PROCYSBI was dorizon may modify a concerning cover specific requirements on By Your Side.	ocused services, including providing logistical and HIPAA authorization that allows me to share protect in behalf of Horizon) to assess insurance coverage for d patient only and is not being made in exchange from the based solely on my professional determination of yor terminate the Program at any time without notice age or reimbursement for any item or service. Ints could result in outreach to the prescriber. Please note that your patient will not benefit fror	
, L		Date				
Written or e-signature only;	(Dispense as Written)			(Sı	ubstitution Permitted)	



PROCYSBI® (CYSTEAMINE BITARTRATE) DELAYED-RELEASE CAPSULES AND DELAYED-RELEASE ORAL GRANULES PATIENT ENROLLMENT FORM

HORIZON.

Phone: 1-855-888-4004 Fax: 1-877-773-9411 PROCYSBI.com

 $Please \ fax\ completed\ form\ to\ 1-877-773-9411, or\ email\ it\ to\ \underline{PROHBYS@horizontherapeutics.com}.$

Patient Consent for Patient Information, Enrolling in Services, and Accessing Financial Support (Referred to as "Patient Authorization")

I hereby authorize my healthcare providers, my health insurance carriers, and my pharmacies to use and disclose my individually identifiable health information, including my medical records, insurance coverage information, and my name, address and telephone number to Horizon Therapeutics USA, Inc. and its affiliates and their respective agents and representatives (collectively, "Horizon"), including third parties authorized by Horizon to administer drug support and to dispense drugs (collectively, "Horizon By Your Side") for the following purposes: (1) to establish eligibility for benefits; (2) to communicate with my healthcare providers and me about my treatment or condition and related products; (3) to facilitate the provision of products, supplies, or services by a third party including, but not limited to, specialty pharmacies; (4) to register me in any applicable product registration program required for my treatment; (5) to enroll me in eligible patient support programs offered by Horizon By Your Side and/or Horizon, including nursing or patient access support services (government-reimbursed programs may not be eligible for all support services offered; please contact Horizon By Your Side for determination); and (6) to send me marketing information or offer me products and services related to my treatment or condition (or other products or services in which I might be interested) and to contact me occasionally to obtain my feedback (for market research purposes only) about my treatment, my condition, or my experience with Horizon and/or Horizon By Your Side otherwise as required or permitted by law. Further, I appoint the Program, on my behalf, to proceed with Program services and to convey this prescription to the dispensing pharmacy, to the extent permitted under state law. I understand the pharmacies may receive a fee from Horizon in exchange for (1) providing me with certain materials and information described above, and (2) using or disclosing certain health information pursuant to this Authorization

I understand that Horizon, as well as my healthcare providers, cannot require me, as a condition of having access to medications, prescription drugs, treatment, or other care, to sign this Authorization. I understand that I am entitled to a copy of this Authorization.

I understand that information disclosed pursuant to this Authorization in some cases may be redisclosed by the recipient and no longer protected by HIPAA or other privacy laws. But Horizon has agreed to use and disclose my information only for purposes of operating the Program. I understand that I may cancel this Authorization at any time by mailing a signed letter requesting such cancellation to Horizon By Your Side, 1 Horizon Way, Deerfield, IL 60015, but that this cancellation will not apply to any information used or disclosed by my healthcare providers and/or health insurance carriers based on this Authorization before they are notified that I have cancelled it. Unless required by state law, this Authorization is valid for whichever is greater: (a) the duration of remaining on this treatment or (b) 10 years from the date signed below. A photocopy of this Authorization will be treated in the same manner as the original.

Date:						
Patient's Printed Name:						
Patient's/Legally Authorized Representative's Signature:						
Legally Authorized Representative's Printed Name (if required):						
Patient's/Legally Authorized Representative's Home Address:						
Street Address:						
City: State: Zip Code:						
Patient's/Legally Authorized Representative's Telephone:						
Patient's/Legally Authorized Representative's Email Address:						
Legally Authorized Representative's Relationship to Patient: Spouse Parent/Legal Guardian Representative per Power of Attorney						
Is there someone else with whom we may discuss your protected health information? \square No \square Yes						
Name:						
Name:						
Relationship to you:						



PROCYSBI DOSING INFORMATION FOR HEALTHCARE PRESCRIBERS



PROCYSBI is available as¹: 25 mg: 60 delayed-release capsules/bottle 75 mg: 60 delayed-release packets/box 300 mg: 120 delayed-release packets/box

Patients starting PROCYSBI who are cysteamine naïve¹

- · Initiate cysteamine treatment immediately after diagnosis of nephropathic cystinosis
- Patients should be started on PROCYSBI at a fraction (1/6 to 1/4) of the maintenance dosage and gradually titrated up to the maintenance dosage over 4 to 6 weeks
 - o Patients 1 year to less than 6 years: Increase the dosage in 10% increments to the maintenance dosage, while monitoring white blood cell (WBC) cystine concentrations. Allow a minimum of 2 weeks between dosage adjustments. If a patient achieves the therapeutic target WBC cystine concentration at a dosage below the recommended weight-based maintenance dosage, then stop dosage escalation and use the dosage as the patient's maintenance dosage
 - o Patients 6 years of age and older: Gradually increase the dosage over 4 to 6 weeks until the maintenance dosage is achieved
- The maintenance dosage after initial dose escalation is 1.3 g/m² of body surface area per day divided into 2 doses given every 12 hours. The table below shows the recommended starting and maintenance dosages of PROCYSBI, converted from body surface area to body weight

Patients converting to PROCYSBI from immediate-release (IR) cysteamine (CYSTAGON)1

• When switching patients from IR cysteamine bitartrate to PROCYSBI, the starting total daily dose of PROCYSBI is equal to the previous total daily dose of IR cysteamine bitartrate. Divide the total daily dose by 2 and administer every 12 hours

Starting and Maintenance Dosage of PROCYSBI by Body Weight in Cysteamine-Naïve Patients 1 Year of Age and Older (Dosage Rounded Using Available Capsule and Packet Strengths)						
Weight in kg		ge in mg Every 12 Hours, Maintenance Dosage	Maintenance PROCYSBI Dosage			
	1/6 of dosage	1/4 of dosage	in mg Every 12 Hours*			
5 or less	25	50	200			
6 to 10	50	75	300			
11 to 15	75	100	400			
16 to 20	100	125	500			
21 to 25	100	150	600			
26 to 30	125	175	700			
31 to 40	125	200	800			
41 to 50	150	225	900			
51 and greater	175	250	1000			

 $^{^*}$ Higher dosages may be required to achieve target therapeutic WBC cystine concentration.

Monitoring dosage¹

- If a patient's precise calculated dosage cannot be obtained, round to the nearest 25 mg for capsules or 75 mg for packets. Only use whole capsules and packets
- After maintenance dosage of PROCYSBI has been achieved, measure the WBC cystine concentration and titrate the PROCYSBI dosage as needed to achieve target WBC cystine concentrations
- If a dosage adjustment is necessary, increase the dosage by 10%. For patients 1 year to less than 6 years of age, allow a minimum of 2 weeks between dose increments. The maximum dosage of PROCYSBI is 1.95 g/m² per day

If tolerability issues occur with PROCYSBI¹

- If adverse reactions occur, decrease the PROCYSBI dosage and then gradually increase to the maintenance dosage
- For cysteamine-naïve patients who have initial intolerance, temporarily discontinue PROCYSBI and then restart at a lower dosage and gradually increase to the maintenance dosage

Please click here for the Full Prescribing Information for complete dosing and administration instructions.

Adherence to cystine-depleting therapy is critical for optimal cystine control^{2,3}

Patients/caregivers should be urged to take PROCYSBI consistently according to the dosing schedule recommended in the prescribing information¹

References: 1. PROCYSBI (cysteamine bitartrate) delayed-release capsules and delayed-release oral granules [prescribing information] Horizon. 2. Gahl WA, Thoene JG, Schneider JA. Cystinosis. N Engl J Med. 2002;347(2):111-121. 3. Brodin-Sartorius A, Tête M-J, Niaudet P, et al. Cysteamine therapy delays the progression of nephropathic cystinosis in late adolescents and adults. Kidney Int. 2012;81(2):179-189.



INDICATION and IMPORTANT SAFETY INFORMATION



INDICATION

PROCYSBI (cysteamine bitartrate) delayed-release capsules and delayed-release oral granules is a cystine-depleting agent indicated for the treatment of nephropathic cystinosis in adults and pediatric patients 1 year of age and older.

IMPORTANT SAFETY INFORMATION

CONTRAINDICATIONS

 Patients with serious hypersensitivity reaction, including anaphylaxis to penicillamine or cysteamine.

WARNINGS AND PRECAUTIONS

- Ehlers-Danlos-like Syndrome: Skin and bone lesions that resemble clinical findings for Ehlers-Danlos-like syndrome have been reported in patients treated with high doses of immediate-release cysteamine bitartrate or other cysteamine salts. Monitor patients for development of skin or bone lesions and reduce PROCYSBI dosing if patients develop these lesions.
- Skin Rash: Severe skin rashes such as erythema multiforme bullosa or toxic epidermal necrolysis have been reported in patients receiving immediate release cysteamine bitartrate. Discontinue use if severe skin rash occurs.
- Gastrointestinal (GI) Ulcers and Bleeding: GI ulceration and bleeding
 have been reported in patients receiving immediate-release cysteamine
 bitartrate. Monitor for GI symptoms and consider decreasing the dose if
 severe symptoms occur.
- Fibrosing Colonopathy: Fibrosing colonopathy has been reported with
 postmarketing use of PROCYSBI. Evaluate patients with severe, persistent,
 and/or worsening abdominal symptoms for fibrosing colonopathy. If the
 diagnosis is confirmed, permanently discontinue PROCYSBI and switch to
 immediate-release cysteamine bitartrate capsules.
- Central Nervous System (CNS) Symptoms: CNS symptoms such as seizures, lethargy, somnolence, depression, and encephalopathy have been associated with immediate-release cysteamine. Monitor for CNS symptoms; interrupt or reduce the dose for severe symptoms or those that persist or progress.
- Leukopenia and/or Elevated Alkaline Phosphatase Levels: Cysteamine
 has been associated with reversible leukopenia and elevated alkaline
 phosphatase levels. Monitor white blood cell counts and alkaline
 phosphatase levels; decrease or discontinue the dose until values revert
 to normal.
- Benign Intracranial Hypertension: Benign intracranial hypertension
 (pseudotumor cerebri; PTC) and/or papilledema has been reported in
 patients receiving immediate-release cysteamine bitartrate treatment.
 Monitor for signs and symptoms of PTC; interrupt or reduce the dose for
 signs/symptoms that persist, or discontinue if diagnosis is confirmed.

ADVERSE REACTIONS

The most common adverse reactions reported in PROCYSBI clinical trials (≥ 5%): were:

- Patients 2 years of age and older previously treated with cysteamine: vomiting, nausea, abdominal pain, headache, conjunctivitis, influenza, gastroenteritis, nasopharyngitis, dehydration, ear infection, upper respiratory tract infection, fatigue, arthralgia, cough, and pain in extremity.
- Patients 1 year of age and older naïve to cysteamine treatment: vomiting, gastroenteritis/viral gastroenteritis, diarrhea, breath odor, nausea, electrolyte imbalance, headache.

DRUG INTERACTIONS

- Drugs that increase gastric pH may alter the pharmacokinetics of cysteamine due to the premature release of cysteamine from PROCYSBI and increase WBC cystine concentration. Monitor WBC cystine concentration with concomitant use.
- Consumption of alcohol with PROCYSBI may increase the rate of cysteamine release and/or adversely alter the pharmacokinetic properties, as well as the effectiveness and safety of PROCYSBI.
- PROCYSBI can be administered with electrolyte (except bicarbonate) and mineral replacements necessary for management of Fanconi Syndrome as well as vitamin D and thyroid hormone.

USE IN SPECIFIC POPULATIONS

 Lactation: Because of the potential risk for serious adverse reactions in breastfed children from cysteamine, breastfeeding is not recommended during treatment with PROCYSBI.

Please see Full Prescribing Information.