## **AvMed**

## PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST\*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid)</u> on this request. All other information may be filled in by office staff; <u>fax to 1-305-671-0200</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. <u>If the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

## tiopronin products

<b>Drug Requested:</b> (Check app	plicable drug below)	
□ tiopronin (generic Thiola®	tiopronin delayed- release tablets (generic Thiola® EC)	□ venxxiva (generic Thiola®)
MEMBER & PRESCRIB	ER INFORMATION: Authorizat	ion may be delayed if incomplete.
Member Name:		
	ber AvMed #: Date of Birth:	
Prescriber Name:		
Prescriber Signature:	escriber Signature: Date:	
Office Contact Name:		
Phone Number: Fax Number:		
NPI #:		
DRUG INFORMATION:	Authorization may be delayed if incom	nplete.
Drug Name/Form/Strength:		
Dosing Schedule:	Length of	Therapy:
Diagnosis:	ICD Code, if applicable:	
Weight (if applicable):	Date	weight obtained:
	Check below all that apply. All criteria ocumentation, including lab results, diaged.	
<b>Initial Authorization: 6 m</b>	onths	
☐ Provider requesting this m with homozygous cystinu	nedication is a nephrologist or has experiria	rience in treating/monitoring members

<u>AND</u>

(Continued on next page)

	Member has a confirmed diagnosis of homozygous cystinuria (documentation recording family history, history of nephrolithiasis, kidney stone collection analysis, and metabolic testing/24-hour urinalysis <a href="MUST">MUST</a> accompany request)		
	AND		
	Before any treatment for cystinuria, the urine cystine levels have been measured to be greater than 500 mg/day (laboratory results MUST be attached to request)		
	Laboratory Results: Date of test:		
	AND		
	Prevention of recurrent cystine stones in this member has <u>NOT</u> been achieved with increased fluid intake, restriction of sodium/protein intake, and urinary alkalinization (ALL THERAPY TRIALS/FAILURES MUST BE RECORDED, DOCUMENTED AND SUBMITTED WITH THIS REQUEST)		
	AND		
	A baseline urinary protein level has been measured, and there are <b>NOT</b> signs of proteinuria		
	Laboratory Results: Date of test:		
	AND		
	A lower dose will be initiated for members who have experienced severe toxicity with D-Penicillamine		
	AND		
	<b>FOR PEDIATRIC PATIENTS:</b> Current weight is $\geq 20 \text{ kg}$		
	Current weight measurement: Date of measurement:		
	(NOTE: tiopronin (Thiola) or tiopronin DR (Thiola EC) will $\underline{NOT}$ be approved for members less than 20 kg, or for doses greater than 50 mg/kg)		
suppo	uthorization: 12 months. Check below all that apply. All criteria must be met for approval. To ort each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be ded or request may be denied.		
and th	E: Renal function, 24-hour urinary protein and urinalysis should have been measured at baseline ten every 3 to 6 months, and urinary cystine level measured 1 month after initiating treatment and every 3 months thereafter		
	Member does <u>NOT</u> have signs of proteinuria (Provide the last interval of urinalysis measuring urinary protein – laboratory results <u>MUST</u> be attached to request)		
	Laboratory Results: Date of test:		
	AND		
(Continued on next page)			

	Provide the last interval of urinalysis measuring urinary cystine levels (laboratory results <u>MUST</u> be attached to request)		
	Laboratory Results:	Date of test:	
	NOTE: Maintenance dose should be ad	djusted to reduce urinary cystine concentration < 250 mg/L	
	AND		
	Improvement/reduction in cystine crystall MUST be attached to request)	uria has been observed and documented (follow up chart notes	
Medication being provided by Specialty Pharmacy – Proprium Rx			

Not all drugs may be covered under every Plan

If a drug is non-formulary on a Plan, documentation of medical necessity will be required.

\*\*Use of samples to initiate therapy does not meet step edit/ preauthorization criteria. \*\*

\*Previous therapies will be verified through pha rmacy paid claims or submitted chart notes. \*