

PENICILLAMINE

Generic	Brand	HICL	GCN	Medi-Span	Exception/Other
PENICILLAMINE	CUPRIMINE,		7091	GPI-14	
	PENICILLAMINE			(99200030000110)	
PENICILLAMINE	DEPEN,		7100	GPI-14	
	PENICILLAMINE			(99200030000305)	
PENICILLAMINE	D-PENAMINE		7101	GPI-14	
				(99200030000302)	

GUIDELINES FOR USE

INITIAL CRITERIA (NOTE: FOR RENEWAL CRITERIA SEE BELOW)

1. Is the request for D-Penamine and the patient has an active prior authorization approval for Depen? [Note: D-Penamine is temporarily available to address a critical drug shortage of Depen. Patients previously approved for Depen will be allowed access without additional criteria during this shortage.]

If yes, approve D-Penamine for 12 months by GPID or GPI-14 for the requested indication as follows:

• Wilson's Disease: #16 per day.

• Active Rheumatoid Arthritis: #12 per day.

• Cystinuria: #32 per day.

If no, continue to #2.

- 2. Does the patient have a diagnosis of Wilson's disease and meet **ALL** of the following criteria?
 - Therapy is prescribed by or in consultation with a hepatologist or gastroenterologist
 - The patient has a Leipzig score of 4 or greater
 - The patient is willing to follow a diet avoiding high copper foods (e.g., shellfish, nuts, chocolate, mushrooms, organ meat)

If yes, continue to #3. If no, continue to #5.

3. Is the request for Depen or D-Penamine?

If yes, approve for 12 months by GPID or GPI-14 for the requested drug as follows:

Depen: #8 per day.

• D-Penamine: #16 per day.

If no, continue to #4.

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INITIAL CRITERIA (CONTINUED)

4. Is the request for Cuprimine and the patient had a trial of or contraindication to Depen (penicillamine) or D-Penamine (penicillamine)?

If yes, approve Cuprimine for 12 months by GPID or GPI-14 with a quantity limit of #8 per day.

If no, do not approve.

DENIAL TEXT: See the initial denial text at the end of the guideline.

- 5. Does the patient have a diagnosis of cystinuria and meet ALL of the following criteria?
 - Therapy is prescribed by or in consultation with a nephrologist
 - The patient has a daily cystine output that is greater than 300mg per 24 hours following urine cystine excretion testing
 - The patient has failed to respond to an adequate trial of or has a contraindication to conventional therapy which includes ALL of the following: increased fluid intake, modest reductions in sodium and protein intake, and urinary alkalinization

If yes, continue to #6. If no, continue to #9.

- 6. Does the patient have nephrolithiasis and meet **ONE** of the following criteria?
 - The patient's stone analysis shows a presence of cystine
 - The patient's urinalysis shows pathognomonic hexagonal cystine crystals
 - The patient has a family history of cystinuria AND a positive cyanide-nitroprusside screening

If yes, continue to #7. If no, do not approve.

DENIAL TEXT: See the initial denial text at the end of the guideline.

7. Is the request for Depen or D-Penamine?

If yes, approve for 12 months by GPID or GPI-14 for the requested drug as follows:

- Depen: #16 per day.
- D-Penamine: #32 per day.

If no, continue to #8.

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INITIAL CRITERIA (CONTINUED)

8. Is the request for Cuprimine and has the patient had a trial of or contraindication to Depen (penicillamine) or D-Penamine (penicillamine) **AND** Thiola (tiopronin)?

If yes, approve Cuprimine for 12 months by GPID or GPI-14 with a quantity of #16 per day. If no, do not approve.

DENIAL TEXT: See the initial denial text at the end of the guideline.

- 9. Does the patient have a diagnosis of active rheumatoid arthritis and meet **ALL** of the following criteria?
 - Therapy is prescribed by or in consultation with a rheumatologist
 - The patient does not have a history or other evidence of renal insufficiency
 - The patient has failed to respond to an adequate trial of at least 3 months of conventional therapy including at least ONE of the following DMARD (disease-modifying antirheumatic drug) agents: methotrexate dose greater than or equal to 20mg per week or maximally tolerated dose, leflunomide, hydroxychloroquine, or sulfasalazine

If yes, continue to #10. If no, do not approve.

DENIAL TEXT: See the initial denial text at the end of the guideline.

10. Is the request for Depen or D-Penamine?

If yes, approve for 12 months by GPID or GPI-14 for the requested drug as follows:

- Depen: #6 per day.
- D-Penamine: #12 per day.

If no, continue to #11.

11. Is the request for Cuprimine and has the patient had a trial of or contraindication to Depen (penicillamine) or D-Penamine (penicillamine)?

If yes, approve Cuprimine for 12 months by GPID or GPI-14 with a quantity of #6 per day. If no, do not approve.

DENIAL TEXT: See the initial denial text at the end of the guideline.

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INITIAL CRITERIA (CONTINUED)

INITIAL DENIAL TEXT: *Some terms are already pre-defined in parenthesis. Please use these definitions if the particular text you need to use does not already have definition(s) in it.

Our guideline named **PENICILLAMINE** (Cuprimine, Depen, D-Penamine) requires the following rule(s) be met for approval:

- A. You have ONE of the following diagnoses:
 - 1. Wilson's disease (a genetic disorder that leads to copper accumulation in the organs)
 - 2. Cystinuria (a type of genetic metabolic disorder)
 - 3. Active rheumatoid arthritis (a type of joint condition)

B. If you have Wilson's disease, approval also requires:

- 1. Therapy is prescribed by or in consultation with a hepatologist (a type of liver doctor) or gastroenterologist (a type of digestive system doctor)
- 2. You have a Leipzig score of 4 or greater (a type of diagnostic score)
- 3. You are willing to follow a diet avoiding high copper foods (such as shellfish, nuts, chocolate, mushrooms, organ meat)
- 4. If you are requesting Cuprimine, you had a trial of or have a contraindication (harmful for) to Depen (penicillamine) or D-Penamine (penicillamine)

C. If you have cystinuria, approval also requires:

- 1. Therapy is prescribed by or in consultation with a nephrologist (kidney doctor)
- 2. You have a daily cystine output greater than 300mg per 24 hours after urine cystine excretion testing
- 3. You have failed to respond to an adequate trial of or has a contraindication (harmful for) to conventional therapy which includes ALL of the following:
 - a. Increased fluid intake
 - b. Modest reductions in sodium and protein intake
 - c. Urinary alkalinization (a process that makes urine basic)
- 4. You have nephrolithiasis (kidney stones) and ONE of the following:
 - a. Your kidney stone analysis shows that there is a presence of cystine (an amino acid)
 - b. Your urine analysis shows that there are hexagonal cystine crystals in your urine that are pathognomonic (signs relating to the disease)
 - c. You have a family history of cystinuria and positive test results in the cyanidenitroprusside screen (a test to determine the amount of cysteine in your body)
- 5. If you are requesting Cuprimine, you had a trial of or have a contraindication (harmful for) to Depen (penicillamine) or D-Penamine (penicillamine) AND Thiola (tiopronin)

(Initial denial text continued on next page)

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INITIAL CRITERIA (CONTINUED)

- D. If you have active rheumatoid arthritis, approval requires:
 - 1. Therapy is prescribed by or in consultation with a rheumatologist (a type of immune system doctor)
 - 2. You do not have a history or other evidence of renal insufficiency (kidney problems)
 - 3. You have failed to respond to an adequate trial of at least 3 months of conventional therapy including at least ONE of the following DMARD (disease-modifying antirheumatic drug) agents: methotrexate dose greater than or equal to 20mg per week or maximally tolerated dose, leflunomide, hydroxychloroquine, or sulfasalazine
 - 4. If you are requesting Cuprimine, you had a trial of or have a contraindication (harmful for) to Depen (penicillamine) or D-Penamine (penicillamine)
- E. If you have an active prior authorization approval for Depen, D-Penamine will be approved without meeting additional criteria during the period of Depen shortage.

Your doctor told us [INSERT PT SPECIFIC INFO PROVIDED]. We do not have information showing you [INSERT UNMET CRITERIA]. This is why your request is denied. Please work with your doctor to use a different medication or get us more information if it will allow us to approve this request.

RENEWAL CRITERIA

- 1. Does the patient have a diagnosis of Wilson's disease **AND** meet the following criterion?
 - The patient has achieved a free serum copper of less than 10 mcg/dL

If yes, approve for lifetime by GPID or GPI-14 for the requested drug as follows:

- Depen: #8 per day.
- Cuprimine: #8 per day.
- D-Penamine: #16 per day.

If no, continue to #2.

- 2. Does the patient have a diagnosis of cystinuria AND meet the following criterion?
 - The patient has achieved a cystine excretion of less than 200 mg/day

If yes, approve for lifetime by GPID or GPI-14 for the requested drug as follows:

- Depen: #16 per day.
- D-Penamine: #32 per day.
- Cuprimine: #16 per day.

If no, continue to #3.

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RENEWAL CRITERIA (CONTINUED)

- 3. Does the patient have a diagnosis of active rheumatoid arthritis and meet **ALL** of the following criteria?
 - The patient does not have a history of or other evidence of renal insufficiency
 - The patient has experienced or maintained improvement in tender joint count or swollen joint count while on therapy compared to baseline

If yes, approve for lifetime by GPID or GPI-14 for the requested drug as follows:

- Depen: #6 per day.
- D-Penamine: #12 per day.
- Cuprimine: #6 per day.

If no, do not approve.

RENEWAL DENIAL TEXT: *Some terms are already pre-defined in parenthesis. Please use these definitions if the particular text you need to use does not already have definition(s) in it.

Our guideline named **PENICILLAMINE** (Cuprimine, Depen, D-Penamine) requires the following rule(s) be met for renewal:

- A. You have ONE of the following diagnoses:
 - 1. Wilson's disease (a genetic disorder that leads to copper accumulation in the organs)
 - 2. Cystinuria (a type of genetic metabolic disorder)
 - 3. Active rheumatoid arthritis (a type of joint condition)
- B. If you have Wilson's disease, approval also requires:
 - 1. You have achieved a free serum copper of less than 10 mcg/dLl
- C. If you have cystinuria, approval also requires:
 - 1. You have achieved a cystine excretion of less than 200 mg/day
- D. If you have active rheumatoid arthritis, approval also requires:
 - 1. You do not have a history of or other evidence of renal insufficiency (kidney problems)
 - 2. You have experienced or maintained improvement in tender joint count or swollen joint count while on therapy compared to baseline

Your doctor told us [INSERT PT SPECIFIC INFO PROVIDED]. We do not have information showing you [INSERT UNMET CRITERIA]. This is why your request is denied. Please work with your doctor to use a different medication or get us more information if it will allow us to approve this request.

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RATIONALE

For further information, please refer to the Prescribing Information and/or Drug Monograph for Cuprimine/Depen/Thiola EC.

REFERENCES

- Cuprimine [Prescribing Information]. Bridgewater, NJ: Bausch Health Companies Inc.; October 2020
- Thiola [Prescribing Information]. San Antonio, TX: Mission Pharmacal; March 2021.
- Depen [Prescribing Information]. Somerset, NJ: Meda Pharmaceuticals; January 2019.
- FDA Website: Penicillamine (Depen) Titratable Tablets Drug Shortage. Available at:
 https://www.accessdata.fda.gov/scripts/drugshortages/dsp_ActiveIngredientDetails.cfm?Al=Penicillamine%20(Depen)%20Titratable%20Tablets&st=c. Accessed on January 21, 2019

Library	Commercial	NSA
Yes	Yes	No

Part D Effective: N/A Created: 05/16

Commercial Effective: 05/08/23 Client Approval: 04/23 P&T Approval: 10/22

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