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HIGHLIGHTS OF PRESCRIBING INFORMATION These highlights do not include all the information needed to use ELCYS <sup>®</sup> safely and effectively. See full prescribing information for ELCYS <sup>®</sup> . ELCYS <sup>®</sup> (cysteine hydrochloride injection), for intravenous use Initial U.S. Approval: 1971				
<b>INDICATIONS AND USAGE</b> ELCYS <sup>®</sup> is a sulfur-containing amino acid indicated to meet the nutritional requirements of newborn infants requiring total parenteral nutrition (TPN); and of adult and pediatric patients with severe liver disease who may have impaired enzymatic processes and require TPN. It can also be added to amino acid solutions to provide a more complete profile of amino acids for protein synthesis. (1)				
<ul> <li>ELCYS<sup>®</sup> is for admixing use only. Not for direct intravenous infusion. (2.1)</li> <li>PHARMACY BULK PACKAGE: Dispense single-doses to many patients in a pharmacy admixture program. Use within 4 hours of puncture. (2.1)</li> <li>See full prescribing information for information on preparation, administration, instructions for use, dosing considerations, including the recommended dosage in pediatric patients and adults. (2.1, 2.2, 2.3, 2.4, 2.5)</li> </ul>				
DOSAGE FORMS AND STRENGTHS				
Injection				
• Pharmacy Bulk Package: 2500 mg/50 mL (50 mg/mL) cysteine hydrochloride, USP in a 50 mL vial. (3)				
CONTRAINDICATIONS				
<ul> <li>Hypersensitivity to one or more amino acids (4)</li> <li>Inborn errors of amino acid metabolism (4)</li> <li>Pulmonary edema or acidosis due to low cardiac output (4)</li> </ul>				
<ul> <li>Pulmonary Embolism due to Pulmonary Vascular Precipitates: If signs of pulmonary distress occur, stop the infusion and initiate a medical evaluation. (5.1)</li> <li>Vein Damage and Thrombosis: Solutions with osmolarity of 900 mOsm/L or more must be infused through a central catheter. (2.1, 5.2)</li> <li>Increased blood urea nitrogen (BUN): Monitor laboratory parameters and discontinue if exceeds normal postprandial limits and continues to increase. (5.3)</li> <li>Acid-Base Imbalance: Monitor laboratory parameters and supplement with electrolytes as needed. (5.4)</li> <li>Hepatobiliary Disorders: Monitor liver function parameters and ammonia levels. (5.5)</li> <li>Hyperammonemia: Neurocognitive delay possible in infants; monitor blood ammonia levels. (5.6, 8.4)</li> <li>Aluminium Toxicity: Increased risk in patients with renal impairment, including preterm infants. (5.7, 8.4)</li> <li>Monitoring and Laboratory Tests: Monitor fluid and electrolytes, serum osmolarity, blood glucose, kidney and liver function, blood count and coagulation parameters throughout treatment. (5.8)</li> </ul>				
ADVERSE REACTIONS Most common adverse reactions are local reactions (warm sensation, erythema, phlebitis and thrombosis at the infusion site), generalized flushing, fever and nausea (6). To report SUSPECTED ADVERSE REACTIONS, contact Exela Pharma Sciences, LLC or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch. See 17 for PATIENT COUNSELING INFORMATION.				

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\* Sections or subsections omitted from the full prescribing information are not listed.

# FULL PRESCRIBING INFORMATION

# **1 INDICATIONS AND USAGE**

ELCYS<sup>®</sup> is indicated for use as an additive to amino acid solutions to meet the nutritional requirements of newborn infants requiring total parenteral nutrition (TPN); and of adult and pediatric patients with severe liver disease who may have impaired enzymatic

processes and require TPN. It can also be added to amino acid solutions to provide a more complete profile of amino acids for protein synthesis.

# 2 DOSAGE AND ADMINISTRATION

#### 2.1 Important Administration Information

Prior to administration, ELCYS® *must be diluted and used as an admixture* in PN solutions. ELCYS® is not for direct intravenous infusion.

The resulting solution is for intravenous infusion into a central or peripheral vein. The choice of a central or peripheral venous route should depend on the osmolarity of the final infusate. Solutions with osmolarity of 900 mOsm/L or greater must be infused through a central catheter [see Warnings and Precautions (5.2)].

ELCYS<sup>®</sup> is supplied as a pharmacy bulk package, and consists of one pharmacy bulk vial which must be diluted prior to intravenous administration *[see Dosage andAdministration (2.3)]*. As a pharmacy bulk vial, ELCYS<sup>®</sup> is intended for dispensing of single doses to many patients. Dispensing from a pharmacy bulk vial should be completed within 4 hours after the vial is penetrated. Do not administer ELCYS<sup>®</sup> as a direct, undiluted intravenous injection.

#### 2.2 Preparation and Administration Instructions

Prior to administration, ELCYS® *must be diluted and used as an admixture* in PN solutions. ELCYS® is not for direct intravenous infusion.

- ELCYS<sup>®</sup> is to be prepared only in a suitable work area such as a laminar flow hood (or an equivalent clean air compounding area). The key factor in the preparation is careful aseptic technique to avoid inadvertent touch contamination during mixing of solutions and addition of other nutrients.
- ELCYS<sup>®</sup> is for addition to amino acid solutions prior to further admixing with dextrose injection using a PN container.
- Use a dedicated line for PN solutions.
- Intravenous lipid emulsions can be infused concurrently into the same vein as ELCYS<sup>®</sup> containing amino acid and dextrose solutions by a Y-connector located near the infusion site; flow rates of each solution should be controlled separately by infusion pumps.
- For administration without lipid emulsion, use a 0.22 micron in-line filter.
- To prevent air embolism, use a non-vented infusion set or close the vent on a vented set, avoid multiple connections, do not connect flexible containers in series, fully evacuate residual gas in the container prior to administration, do not pressurize the flexible container to increase flow rates, and if administration is controlled by a pumping device, turn off pump before the container runs dry.
- If infused with lipid emulsion, do not use administration sets and lines that contain di-2-ethylhexyl phthalate (DEHP). Administration sets that contain polyvinyl chloride (PVC) components have DEHP as a plasticizer.
- Visually inspect the diluted PN solution containing ELCYS<sup>®</sup> for particulate matter before admixing, after admixing, and prior to administration. The solution should be clear and there should be no precipitates. A slight yellow color does not alter the quality and efficacy of this product.

#### 2.3 Preparation Instructions for Admixing Using a Parenteral Nutrition (PN) Container

- Remove ELCYS<sup>®</sup> vial from the carton and inspect for particulate matter.
- Transfer the required amount of ELCYS<sup>®</sup> to an amino acid solution using strict aseptic techniques to avoid microbial contamination.
- The amino acid solution containing ELCYS<sup>®</sup> can then be used to prepare admixtures in the PN container using strict aseptic techniques.
- Amino acids solution containing ELCYS<sup>®</sup> may be mixed with dextrose injection. The following proper mixing sequence must be followed to minimize pH related problems:
- Transfer dextrose injection to the parental nutrition pooling container
- Transfer phosphate salt
- Transfer ELCYS<sup>®</sup>-containing amino acid solution
- Transfer electrolytes
- Transfer trace elements
- Use gentle agitation during admixing to minimize localized concentration effects; shake containers gently after each addition.
- For automated compounding, refer to Instructions for Use of the applicable compounder.
- Because additives may be incompatible, evaluate all additions to the PN container for compatibility and stability of the resulting preparation. Consult with pharmacist, if available. Questions about compatibility may be directed to Exela Pharma Sciences, LLC. If it is deemed advisable to introduce additives to the PN container, use aseptic technique.
- Inspect the final PN solution containing ELCYS<sup>®</sup> to ensure that precipitates have not formed during mixing or addition on additives. Discard if any precipitates are observed.

#### Stability and Storage

- Use of ELCYS<sup>®</sup> for admixing should be limited to up to 4 hours at room temperature (25°C/77°F) after the container closure has been penetrated. Discard any remaining drug.
- Use PN solution containing ELCYS<sup>®</sup> promptly after mixing. Any storage of the admixture should be under refrigeration and limited to a brief period of time, no longer than 24 hours. After removal from refrigeration, use promptly and complete the infusion within 24 hours. Discard any remaining admixture.
- Protect PN solution from light.

# 2.4 Dosing Considerations

- The dosage of the final PN solution containing ELCYS<sup>®</sup> must be based on the concentrations of all components in the solution and the recommended nutritional requirements [see Dosage and Administration (2.5)]. Consult the prescribing information of all added components to determine the recommended nutritional requirements for dextrose and lipid emulsion, as applicable.
- The dosage of ELCYS<sup>®</sup> should be individualized based on the patient's clinical condition (ability to adequately metabolize amino acids), body weight and nutritional/fluid requirements, as well as additional energy given orally/enterally to

the patient. Prior to initiating parenteral nutrition, the following patient information should be reviewed: review of all medications, gastrointestinal function and laboratory data (such as electrolytes (including magnesium, calcium, and phosphorus), glucose, urea/creatinine, liver panel, complete blood count and triglyceride level (if adding lipid emulsion).

• Prior to administration of PN solution containing ELCYS<sup>®</sup>, correct severe fluid, electrolyte and acid-base disorders.

#### 2.5 Recommended Dosage in Pediatric Patients and Adults

The recommended dosage and volume of ELCYS<sup>®</sup> is shown in Table 1 and is based upon the recommended daily protein (amino acids) requirement. For pediatric patients from birth to less than 12 years of age, the recommended dosage of ELCYS<sup>®</sup> is 22 mg/gram of amino acids. For adults and pediatric patients 12 years of age and older, the recommended dosage of ELCYS<sup>®</sup> is 7 mg/gram of amino acids.

# Table 1. Recommended Daily Dosage of $\mathsf{ELCYS}^{\texttt{R}}$ in Pediatric Patients and Adults

Age	Recommended Protein <sup>a</sup> Requirement (g AA/kg/day) <sup>1</sup>	Recommended Dosage (mg ELCYS/g AA)	Recommended Volume (mL ELCYS/g AA)
Preterm and term infants less than 1 month of age	3 to 4	22	0.44
Pediatric patients 1 month to less than 1 year of age	2 to 3	22	0.44
Pediatric patients 1 year to 11 years of age	1 to 2	22	0.44
Pediatric patients 12 years to 17 years of age	0.8 to 1.5	7	0.14
Adults: Stable Patients	0.8 to 1	7	0.14
Adults: Critically III Patients <sup>b</sup>	1.5 to 2	7	0.14

AA = Amino Acid

a Protein is provided as amino acids (AA)

b Includes patients requiring more than 2 to 3 days in the intensive care unit with organ failure, sepsis or postoperative major surgery. Do not use in patients with conditions that are contraindicated [see Contraindications (4)]

ELCYS<sup>®</sup> contains 50 ma/mL of cvsteine hvdrochloride (equivalent to 34.5 ma/mL of

cysteine). Therefore, the ELCYS<sup>®</sup> dosages in Table 1 provide:

- 15 mg cysteine/gram of amino acids for pediatric patients less than 12 years of age
- 5 mg cysteine/gram of amino acids for adults and pediatric patients 12 years of age and older

## **3 DOSAGE FORMS AND STRENGTHS**

Injection

• Pharmacy Bulk Package: 2500 mg/50mL (50 mg/mL) cysteine hydrochloride, USP in a clear, colorless, sterile solution in a 50 mL vial.

# **4 CONTRAINDICATIONS**

ELCYS<sup>®</sup> is contraindicated in:

- Patients with known hypersensitivity to one or more amino acids.
- Patients with inborn errors of amino acid metabolism due to risk of severe metabolic or neurologic complications.
- Patients with pulmonary edema or acidosis due to low cardiac output.

# **5 WARNINGS AND PRECAUTIONS**

#### 5.1 Pulmonary Embolism due to Pulmonary Vascular Precipitates

Pulmonary vascular precipitates causing pulmonary vascular emboli and pulmonary distress have been reported in patients receiving PN. In some fatal cases, pulmonary embolism occurred as a result of calcium phosphate precipitates. Precipitation following passage through an in-line filter and suspected in vivo precipitate formation has also been reported. If signs of pulmonary distress occur, stop the PN infusion and initiate a medical evaluation. In addition to inspection of the solution [see Dosage and Administration (2.1, 2.2)], the infusion set and catheter should also periodically be checked for precipitates.

# 5.2 Vein Damage and Thrombosis

ELCYS<sup>®</sup> must be diluted and used as an admixture in PN solutions. It is not for direct intravenous infusion. Solutions with an osmolarity of 900 mOsm/L or greater must be infused through a central catheter [see Dosage and Administration (2.1)]. The infusion of hypertonic nutrient injections into a peripheral vein may result in vein irritation, vein damage, and/or thrombosis. The primary complication of peripheral access is venous thrombophlebitis, which manifests as pain, erythema, tenderness or a palpable cord. Remove the catheter as soon as possible, if thrombophlebitis develops.

# 5.3 Increased Blood Urea Nitrogen (BUN)

Intravenous infusion of amino acids may induce a rise in blood urea nitrogen (BUN), especially in patients with impaired hepatic or renal function. Appropriate laboratory tests should be performed periodically and infusion discontinued if BUN levels exceed normal postprandial limits and continue to rise. It should be noted that a modest rise in BUN normally occurs as a result of increased protein intake.

Administration of amino acid solutions in the presence of impaired renal function may augment an increasing BUN, as does any protein dietary component.

#### 5.4 Acid-Base Imbalance

Administration of ELCYS<sup>®</sup> may result in metabolic acidosis in preterm infants.

Administration of amino acid solutions to a patient with hepatic impairment may result in serum amino acid imbalances, metabolic alkalosis, prerenal azotemia, hyperammonemia, stupor and coma.

Frequent clinical evaluation and laboratory determinations are necessary for proper monitoring of acid-base balance during parenteral nutrition therapy. Significant deviations from normal concentrations may require the use of additional electrolyte supplements.

# 5.5 Hepatobiliary Disorders

Hepatobiliary disorders are known to develop in some patients without preexisting liver disease who receive PN, including cholecystitis, cholelithiasis, cholestasis, hepatic steatosis, fibrosis and cirrhosis, possibly leading to hepatic failure. The etiology of these disorders is thought to be multifactorial and may differ between patients.

Monitor liver function parameters and ammonia levels. Patients developing signs of hepatobiliary disorders should be assessed early by a clinician knowledgeable in liver diseases in order to identify possible causative and contributory factors, and possible therapeutic and prophylactic interventions.

# 5.6 Hyperammonemia

Hyperammonemia is of special significance in infants, as it can result in neurocognitive delays. Therefore, it is essential that blood ammonia levels be measured frequently in infants.

Instances of asymptomatic hyperammonemia have been reported in patients without overt liver dysfunction. The mechanisms of this reaction are not clearly defined but may involve genetic defects and immature or subclinically impaired liver function [see Contraindications (4), Use in Specific Populations (8.4)].

# 5.7 Aluminum Toxicity

ELCYS<sup>®</sup> contains aluminum that may be toxic.

Aluminum may reach toxic levels with prolonged parenteral administration in patients with renal impairment. Preterm infants are particularly at risk for aluminum toxicity because their kidneys are immature, and they require large amounts of calcium and phosphate solutions, which also contain aluminum.

Patients with renal impairment, including preterm infants, who receive greater than 4 to 5 mcg/kg/day of parenteral aluminum can accumulate aluminum to levels associated with central nervous system and bone toxicity. Tissue loading may occur at even lower rates of administration.

Exposure to aluminum from ELCYS<sup>®</sup> is not more than 0.21 mcg/kg/day when preterm and term infants less than 1 month of age are administered the recommended maximum dosage of ELCYS<sup>®</sup> (15 mg cysteine/g of amino acids and 4 g of amino acids/kg/day) [see Table 1, Dosage and Administration (2.1)]. When prescribing ELCYS<sup>®</sup> for use in PN containing other small volume parenteral products, the total daily patient exposure to aluminum from the admixture should be considered and maintained at no more than 5 mcg/kg/day [see Use in Specific Populations (8.4)].

## 5.8 Monitoring and Laboratory Tests

Monitor fluid and electrolyte status, serum osmolarity, blood glucose, liver and kidney function, blood count and coagulation parameters throughout treatment [see Dosage and Administration (2.4)].

# **6 ADVERSE REACTIONS**

The following serious adverse reactions are discussed in greater detail in other sections of the prescribing information:

- Pulmonary embolism due to pulmonary vascular precipitates [see Warnings and *Precautions (5.1)*]
- Vein damage and thrombosis [see Warnings and Precautions (5.2)]
- Increased BUN [see Warnings and Precautions (5.3)]
- Acid-base imbalance [see Warnings and Precautions (5.4)]
- Hepatobiliary disorders [see Warnings and Precautions (5.5)]
- Hyperammonemia [see Warnings and Precautions (5.6)]
- Aluminum toxicity [see Warnings and Precautions (5.7)]

Adverse reactions with the use of cysteine hydrochloride injection were identified in clinical studies or postmarketing reports. Because some of these reactions were reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

- Local infusion site reactions, including a warm sensation, erythema, phlebitis and thrombosis at the infusion site
- Generalized flushing, fever and nausea

# **8 USE IN SPECIFIC POPULATIONS**

#### 8.1 Pregnancy

#### <u>Risk Summary</u>

Appropriate administration of ELCYS<sup>®</sup> is not expected to cause major birth defects, miscarriage or adverse maternal or fetal outcomes. Animal reproduction studies have not been conducted with cysteine hydrochloride.

The estimated background risk for major birth defects and miscarriage for the indicated populations is unknown. All pregnancies have a background risk of birth defect, loss or other adverse outcomes. In the U.S. general population, the estimated background risk of major birth defect and miscarriage in the clinically recognized pregnancies is 2 to 4%

and 15 to 20%, respectively.

#### 8.2 Lactation

#### <u>Risk Summary</u>

Data available on the effects of cysteine hydrochloride on infants, either directly or through breastmilk, do not suggest a significant risk of adverse events from exposure. Although there are no data on the presence of cysteine hydrochloride in human or animal milk or the effects on milk production, appropriate administration of ELCYS<sup>®</sup> is not expected to cause harm to a breastfed infant. The development and health benefits of breastfeeding should be considered along with the mother's clinical need for ELCYS<sup>®</sup> and any potential adverse effects on the breastfed infant from ELCYS<sup>®</sup> or from the underlying maternal condition.

#### 8.4 Pediatric Use

ELCYS<sup>®</sup> is approved for use in pediatric patients, from birth to 17 years of age, for use as an additive to amino acid solutions to meet the nutritional requirements of newborn infants, including preterm infants, requiring total parenteral nutrition (TPN) and pediatric patients with severe liver disease who may have impaired enzymatic processes and require TPN. The safety profile for ELCYS<sup>®</sup> use in pediatric patients includes risks of acid-base imbalance and hyperammonemia.

Acid-base imbalance, including metabolic acidosis, may occur with ELCYS<sup>®</sup> administration in preterm infants. Frequent clinical and laboratory assessments are necessary to monitor and manage fluid balance, electrolyte concentrations, and acid-base balance during parenteral nutrition therapy [see Warnings and Precautions (5.4)].

Hyperammonemia is of special significance in infants (birth to two years of age). This reaction appears to be related to a deficiency of the urea cycle amino acids of genetic or product origin. It is essential that blood ammonia be measured frequently in infants [see Warnings and Precautions (5.6)].

Because of immature renal function, preterm infants receiving prolonged parenteral nutrition treatment with ELCYS<sup>®</sup> may be at higher risk of aluminum toxicity [see Warnings and Precautions (5.7)].

# 8.5 Geriatric Use

Clinical studies with ELCYS<sup>®</sup> have not been performed to determine whether patients aged 65 and over respond differently from younger patients.

# 8.6 Renal Impairment

Monitor patients with impaired renal function receiving PN solutions containing the recommended dosage of ELCYS<sup>®</sup> with frequent clinical evaluation and laboratory tests to assess renal function, including serum electrolytes and fluid balance [see Dosage and Administration (2.4), Warnings and Precautions (5.8)].

# 8.7 Hepatic Impairment

Monitor patients with impaired liver function receiving PN solutions containing the recommended dosage of  $ELCYS^{(R)}$  with frequent clinical evaluation and laboratory tests

to assess liver function, such as bilirubin and liver function parameters [see Dosage and Administration (2.4), Warnings and Precautions (5.8)].

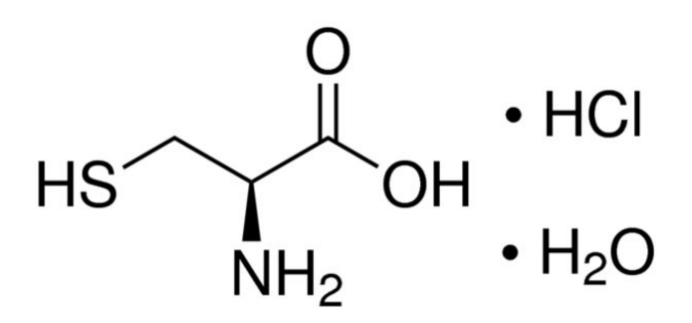
# **10 OVERDOSAGE**

In the event of over hydration or solute overload, re-evaluate the patient and institute appropriate corrective measures [see Warnings and Precautions (5.3, 5.4, 5.5, 5.7, 5.8)].

# **11 DESCRIPTION**

ELCYS<sup>®</sup> (cysteine hydrochloride injection) is a sterile, nonpyrogenic solution for intravenous use. Each 50 mL of ELCYS<sup>®</sup> contains 2500 mg of cysteine hydrochloride, USP (equivalent to 345 mg of cysteine) in water for injection. Sodium hydroxide and/or hydrochloric acid are used as needed to adjust the pH. The pH range is 1.0 to 2.5.

The active ingredient is cysteine hydrochloride. Cysteine is a sulfur-containing amino acid. The chemical name of cysteine hydrochloride is L-cysteine hydrochloride monohydrate and is chemically designated as C<sub>3</sub>H<sub>7</sub>NO<sub>2</sub>S • HCI • H<sub>2</sub>O having a molecular weight of 175.63. Cysteine hydrochloride is a white crystalline powder soluble in water. Cysteine aqueous solution is prone to oxidation when exposed to air, and when mixed with amino acids solutions, cysteine may convert to insoluble cystine which leads to precipitation over time. It has the following structural formula:



ELCYS<sup>®</sup> contains no more than 80 mcg/L of aluminum.

# **12 CLINICAL PHARMACOLOGY**

# 12.1 Mechanism of Action

Endogenous cysteine is synthesized from methionine by the enzyme, cystathionase, via

the trans-sulfuration pathway, and serves as a precursor substrate for both glutathione and taurine. ELCYS<sup>®</sup> provides cysteine to the systemic circulation of patients who require PN and cannot synthesize adequate quantities of cysteine due to insufficient or deficient cystathionase activity.

#### **15 REFERENCES**

1. Ayers P. et al. A.S.P.E.N. Parenteral Nutrition Handbook, 2nd ed. 2014 pg. 123 and 124.

#### **16 HOW SUPPLIED/STORAGE AND HANDLING**

ELCYS<sup>®</sup> is supplied as follows:

Pharmacy Bulk Package: 2,500 mg/50 mL (50 mg/mL) of cysteine hydrochloride, USP is a clear, colorless, sterile and nonpyrogenic solution in 50 mL vials packaged as 1 per carton (NDC 51754-1008-1), 25 cartons per case (NDC 51754-1008-4)

Store at 20°C to 25°C (68°F to 77°F). [See USP Controlled Room Temperature]. Avoid excessive heat. Protect from freezing. If accidentally frozen, discard the vial.

For storage of admixed solution see *Dosage and Administration (2.3)*.

#### **17 PATIENT COUNSELING INFORMATION**

Inform patients, caregivers, or home healthcare providers of the following risks of ELCYS<sup>®</sup>:

- Pulmonary embolism due to pulmonary vascular precipitates [see Warnings and *Precautions (5.1)*]
- Vein damage and thrombosis [see Warnings and Precautions (5.2)]
- Increased BUN [see Warnings and Precautions (5.3)]
- Acid base imbalance [see Warnings and Precautions (5.4)]
- Hepatobiliary disorders [see Warnings and Precautions (5.5)]
- Hyperammonemia [see Warnings and Precautions (5.6)]
- Aluminum toxicity [see Warnings and Precautions (5.7)]
- Monitoring and laboratory tests [see Warnings and Precautions (5.8)]

Manufactured and Distributed by:



Exela Pharma Sciences, LLC Lenoir, NC 28645

#### PACKAGE/LABEL PRINCIPAL DISPLAY PANEL-Vial Label



Rx Only NDC 51754-1008-1

**ELCYS**<sup>®</sup>

(cysteine hydrochloride injection), USP

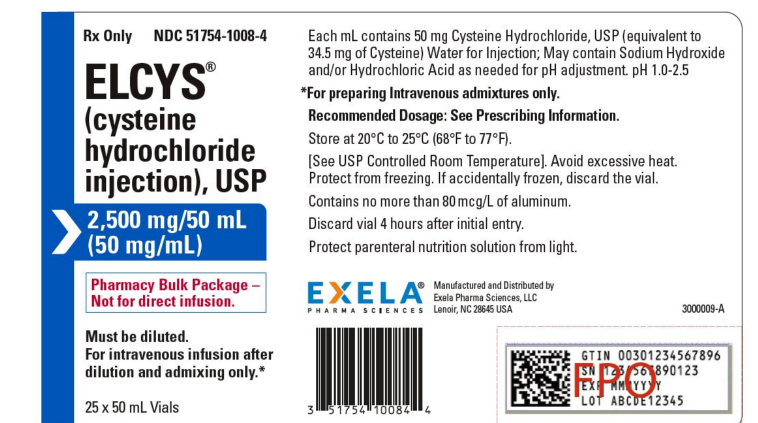
2,500 mg/50 mL (50 mg/mL)

Pharmacy Bulk Package - Not for direct infusion.

Must be diluted. For intravenous infusion after dilution and admixing only.\*

50 mL Vial

# PACKAGE/LABEL PRINCIPAL DISPLAY PANEL-Carton Label



Rx Only NDC 51754-1008-4

**ELCYS**<sup>®</sup>

(cysteine hydrochloride injection), USP

2,500 mg/50 mL (50 mg/mL)

Pharmacy Bulk Package - Not for direct infusion.

Must be diluted. For intravenous infusion after dilution and admixing only.\*

25 x 50 mL Vials

# **ELCYS**

cysteine hydrochloride injection, solution

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	ltem Code (Source)	NDC:51754-1008
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
CYSTEINE HYDROCHLORIDE (UNII: ZT934N0X4W) (CYSTEINE - UNII:K848JZ4886)	CYSTEINE	50 mg in 1 mL	

Inactive Ing	redients				
Ingredient Name Streng					trength
WATER (UNII: 059QF0K00R)					
Other Ingred	dients				
Ingredien	t Kind	Ingredie	nt Name		Quantity
May contain		SODIUM HYDROXIDE (UNII: 55X0	1QC32I)		
May contain		HYDROCHLORIC ACID (UNII: QTT	17582CB)		
	•	Package Description	Marketing Date		Marketing End Date
# Item Code	25 in 1 CA				-
NDC:51754-	25 in 1 CA		Date		-
<ul> <li># Item Code</li> <li>1 NDC:51754- 1008-4</li> </ul>	25 in 1 CAI 50 mL in 1	RTON	Date		-
<ul> <li># Item Code</li> <li>1 NDC:51754- 1008-4</li> <li>1</li> </ul>	25 in 1 CA 50 mL in 1 Product	RTON VIAL; Type 0: Not a Combination	Date		-
<ul> <li># Item Code</li> <li>1 NDC:51754- 1008-4</li> </ul>	25 in 1 CA 50 mL in 1 Product	RTON VIAL; Type 0: Not a Combination	h Marketi		-

Labeler - Exela Pharma Sciences, LLC (831274399)

Establishment			
Name	Address	ID/FEI	Business Operations
Exela Pharma Sciences, LLC		831274399	MANUFACTURE(51754-1008) , PACK(51754-1008) , LABEL(51754- 1008)

Revised: 2/2024

Exela Pharma Sciences, LLC