CLINIMIX- leucine, phenylalanine, lysine, methionine, isoleucine, valine, histidine, threonine, tryptophan, alanine, glycine, arginine, proline, serine, tyrosine, dextrose injection CLINIMIX- leucine, phenylalanine, lysine, methionine, isoleucine, valine, histidine, threonine, tryptophan, alanine, glycine, arginine, proline, serine, tyrosine, dextrose injection CLINIMIX- leucine, phenylalanine, lysine, methionine, isoleucine, valine, histidine, threonine, tryptophan, alanine, glycine, arginine, proline, serine, tyrosine, dextrose injection Baxter Healthcare Corporation

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#### HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use CLINIMIX safely and effectively. See full prescribing information for CLINIMIX.

- Known hypersensitivity to one or more amino acids or dextrose. (4)
- Inborn errors of amino acid metabolism. (4)
- Patients with pulmonary edema or acidosis due to low cardiac output. (4)

------ WARNINGS AND PRECAUTIONS ------

- Pulmonary Embolism due to Pulmonary Vascular Precipitates: if signs of pulmonary distress occur, stop the infusion and initiate a medical evaluation. (5.1)
- Hypersensitivity Reactions: monitor for signs and symptoms and discontinue infusion if reactions occur. (5.2)
- Risk of Infections, Refeeding Complications, and Hyperglycemia or Hyperosmolar Hyperglycemic State: monitor for signs and symptoms; monitor laboratory parameters. (5.3, 5.4, 5.5)
- Vein Damage and Thrombosis: solutions with osmolarity of ≥ 900 mOsm/L must be infused through a central catheter. (2.2, 5.6)
- Hepatobiliary Disorders: monitor liver function parameters and ammonia levels. (5.7)
- Aluminum Toxicity: increased risk in patients with renal impairment, including preterm infants. (5.8, 8.4)
- Parenteral Nutrition Associated Liver Disease: increased risk in patients who receive parenteral nutrition for extended periods of time, especially preterm infants; monitor liver function tests, if abnormalities occur consider discontinuation or dosage reduction. (5.9, 8.4)
- Electrolyte Imbalance and Fluid Overload: patients with cardiac insufficiency or renal impairment may require adjustment of fluid, protein and electrolyte content. (5.10, 8.4)

ADVERSE REACTIONS
Adverse reactions include diuresis, extravasation, glycosuria, hyperglycemia, and hyperosmolar coma. (6)
To report SUSPECTED ADVERSE REACTIONS, contact Baxter Healthcare at 1-866-888-2472 or FDA at 1-800-
FDA-1088 or www.fda.gov/medwatch
USE IN SPECIFIC POPULATIONS
<u>Pediatric Use</u> : increased risk of hypoglycemia/hyperglycemia: monitor serum glucose concentrations. (8.4)
See 17 for PATIENT COUNSELING INFORMATION.

Revised: 10/2018

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#### **FULL PRESCRIBING INFORMATION**

#### 1 INDICATIONS AND USAGE

CLINIMIX is indicated as a source of calories and protein for patients requiring parenteral nutrition when oral or enteral nutrition is not possible, insufficient, or contraindicated. CLINIMIX may be used to treat negative nitrogen balance in patients.

<sup>\*</sup> Sections or subsections omitted from the full prescribing information are not listed.

#### 2 DOSAGE AND ADMINISTRATION

# 2.1 Preparation Prior to Administration

- Tear protective clear overwrap across top at slit and remove solution container. Small amounts of
  moisture may be found on the solution container from water permeating from inside the container.
  The amount of permeated water is insufficient to affect the solution significantly. If larger
  amounts of water are found, the container should be checked for tears or leaks.
- Inspect the container prior to activation. Some opacity of the plastic due to moisture absorption during the sterilization process may be observed. This is normal and does not affect the solution quality or safety. The opacity will diminish gradually. Evaluate the following:
  - O If the outlet or additive port protectors are damaged, detached, or not present, discard container as solution path sterility may be impaired.
  - Oheck to ensure seal between chambers is intact, solutions are contained in separate chambers, and the content of the individual chambers is clear, colorless or slightly yellow. Discard if the seal is broken or if the solution is bright yellow or yellowish brown.
  - O Check for minute leaks by separately squeezing each chamber. If external leaks or leakage between the chambers are found, discard solution as sterility or stability may be impaired.
- Lipids and/or additives can be introduced to the container after opening seal between chambers. Because additives may be incompatible, evaluate all additions to the plastic container for compatibility. Activate chambers of container prior to introduction of additives. Mix thoroughly when additives have been introduced. Supplemental medication may be added with a 19 to 22 gauge needle through the medication port.
- Calcium and phosphate ratios must be considered. Excess addition of calcium and phosphate, especially in the form of mineral salts, may result in the formation of calcium phosphate precipitates [see Warnings and Precautions (5.1)].
- Inspect the container to ensure precipitates have not formed during the mixing or addition of additives. A slight yellow color does not alter the quality and efficacy of this product. If lipid has been added, ensure the emulsion has not separated. Separation of the emulsion can be visibly identified by a yellowish streaking or the accumulation of yellowish droplets in the mixed emulsion. Discard the admixture if any of the above are observed

## 2.2 Important Administration Instructions

- Set the vent to the closed position on a vented intravenous administration set to prevent air embolism.
- Use a dedicated line without any connections to avoid air embolism.
- CLINIMIX is for intravenous infusion only 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 (5.6)].
  - o For central vein infusion only: CLINIMIX 4.25/10, 5/15, 5/20
  - O For central or peripheral vein infusion: CLINIMIX 4.25/5
- The solution should be inspected for precipitates before admixing, after admixing, and again before administration.
- Use a 0.22 micron filter for administration of CLINIMIX. If a lipid is also administered, use a 1.2 micron filter.
- If lipid emulsion is added, 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.

#### 2.3 Instructions for Use

- 1. Open by tearing protective clear overwrap, which includes an oxygen-absorbing sachet, across top at slit and remove solution container. Discard the oxygen-absorbing sachet after removal from the clear overwrap.
- 2. To proceed with activation, the container should be at room temperature. Lay the room temperature container onto a flat surface. Grasp the container firmly on each side of the top of the container (**Figure 1**).
- 3. Starting from the top, using some pressure, slowly roll the container to open seal between chambers as shown in **Figure 2**. Do not pull or rip the seal apart. The seal must be completely opened towards the port side of the container. The upper section of the seal towards the hanger side can remain unbroken.
- 4. Mix the contents thoroughly by inverting the container upside down to ensure a homogenous admixture (**Figure 3**).
- 5. Once the container is mixed, check for leaks.
- 6. Make additions (if prescribed).

Because additives may be incompatible, evaluate all additions to the container for compatibility and stability of the resulting preparation. Consult with pharmacist, if available. Questions about compatibility may be directed to Baxter. If it is deemed advisable to introduce additives, use aseptic technique. For information on adding lipid emulsions *see Dosage and Administration* (2.4).

- a. Prepare medication port.
- b. Using syringe with 19 to 22 gauge needle, puncture resealable medication port and inject.
- C. Mix solution and medication thoroughly (**Figure 3**). For high density medication (high specific gravity), such as potassium chloride, squeeze ports while ports are upright and mix thoroughly.
- 7. Inspect final solution for discoloration and particulate matter. Check for leaks.
- 8. Spike and hang container.
  - d. Suspend container from eyelet support.
  - e. Twist off protector from outlet port at bottom of container (**Figure 4**).
  - f. Attach administration set. Refer to complete directions accompanying set.

For single dose only. Discard unused portion.

# Figures 1-4:





Lay the room temperature container onto a flat surface. Grasp the container firmly with both hands at the top corners.



Starting from the top, using some pressure, slowly roll the container down toward the bottom until the peel seal is broken down the center. You should feel or hear an audible pop as the peel seal dividing the chambers is broken, which allows the components from each chamber to mix.





Mix by turning the container upside-down at least 3 times.

4.



Hang the container. Twist off the protector from the administration outlet. Firmly plug the spike connector.

# **Instructions on Storage**

Storage After Removal of Overwrap:

Once removed from the protective clear overwrap, mixed (peel seal activated) or unmixed (peel seal intact) CLINIMIX solutions may be stored under refrigeration for up to 9 days.

Storage Once any Additive is Added:

Use promptly after mixing. Any storage with additives should be under refrigeration and limited to a brief period of time, less than 24 hours. After removal from refrigeration, use promptly and complete the infusion within 24 hours. Any remaining mixture must be discarded.

# 2.4 Preparation and Addition of Lipid Emulsion

- 1. Prior to adding lipid emulsion, mix amino acid and dextrose injection as shown in **Figures 1-3**.
- 2. Prepare lipid emulsion transfer set following instructions provided.
- 3. Attach transfer set to lipid emulsion container using aseptic technique.
- 4. Twist off protector on the additive port of the container.
- 5. Attach the transfer set to the exposed additive port.
- 6. Open clamp on transfer set.
- 7. After completing transfer, use appropriate plastic clamp or metal ferrule to seal off additive port tube.
- 8. Remove transfer set.
- 9. Mix contents of container thoroughly. Inspect final solution for discoloration and particulate matter. Check for leaks.

*Storage Once Lipids are Added:* 

Use promptly after mixing. Any storage with additives 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. Any mixture remaining must be discarded.

# 2.5 Dosing Considerations

- The dosage of CLINIMIX should be individualized based on the patient's clinical condition (ability to adequately metabolize amino acids and dextrose), body weight and nutritional/fluid requirements, as well as additional energy given orally/enterally to the patient. Prior to initiating CLINIMIX the following patient information should be reviewed: all concomitant 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). Refer to the complete prescribing information of lipid emulsion for dosing information.
- CLINIMIX formulations have varying concentrations of protein and carbohydrate; thus infusion rates to achieve requirements will vary. Protein, caloric, fluid and electrolyte requirements all need to be taken into consideration when determining individual patient dosage needs.
- The dosage selection is based only on the recommended protein requirements. The maximum dextrose infusion rates and calorie and fluid requirements must also be considered when determining the clinically appropriate infusion rate for patients.
- CLINIMIX meets the total nutritional requirements for protein and dextrose in stable patients, and can be individualized to meet specific needs with the addition of nutrients.
- Total daily fluid requirements can be met beyond the volume of amino acids solution by supplementing with non-carbohydrate or carbohydrate-containing electrolyte solutions. In many patients, provision of adequate calories in the form of hypertonic dextrose may require the administration of exogenous insulin to prevent hyperglycemia and glycosuria.
- Prior to administration of CLINIMIX correct severe fluid, electrolyte and acid-base disorders.
- Monitor levels of serum potassium during therapy. It may be necessary to add potassium to the CLINIMIX admixture.
- Lipid emulsion administration should be considered with prolonged use (more than 5 days) of CLINIMIX in order to prevent essential fatty acid deficiency (EFAD). Serum lipids should be monitored for evidence of EFAD in patients maintained on fat-free parenteral nutrition. See prescribing information of lipid emulsion.
- The flow rate should be increased gradually. The flow rate must be adjusted taking into account the dose being administered, the daily volume intake, and the duration of the infusion.

# 2.6 Recommended Dosage in Adults

The recommended daily nutritional requirements for protein and dextrose compared to the amount of nutrition provided by CLINIMIX are shown in **Table 1**.

As indicated on an individual basis, maintenance vitamins, electrolytes, trace elements and other components (including lipids) should be administered as required to prevent deficiencies and complications from developing.

The maximum infusion rates in adult patients are show in Table 2.

In addition to meeting protein needs, the administration rate should be governed, especially during the first few day of therapy, by the patient's tolerance to dextrose. Daily intake of amino acids and dextrose should be increased gradually to the maximum required dose as indicated by frequent determinations of blood glucose levels.

**Table 1: Nutritional Comparison – Adult Patients** 

Recommended	
Nutritional	Recommended CLINIMIX Adult Dosage

	Requirements					
	Stable Patients	Critically Ill Patients*		CLINIMIX 4.25/10	CLINIMIX 5/15	CLINIMIX 5/20
<b>Fluid</b> (mL/kg/day)	30 to 40	Minimum needed to deliver adequate nutrition	19 to 40	19 to 40	16 to 40	16 to 40
<b>Protein</b> <sup>†</sup> (g/kg/day)	0.8 to 1	1.5 to 2	0.8 to 1.7	0.8 to 1.7	0.8 to 2	0.8 to 2
(Nitrogen	(0.13 to)	(0.24 to	(0.13  to  0.27)	(0.13 to 0.27)	(0.13 to 0.32)	(0.13 to 0.32)
g/kg/day)	0.16)	0.32)				
<b>Dextrose</b> (g/kg/day)	≤10	≤5.8	0.95 to 2	1.9 to 4	2.4 to 6	3.2 to 8

<sup>\*</sup> Do not use in patients with conditions that are contraindicated [see *Contraindications (4)*].

**Table 2: Maximum Infusion Rate in Adult Patients:** 

		Maximum Infusion Rates in Adults Patients					
		CLINIMIX 4.25/5	CLINIMIX 4.25/10	CLINIMIX 5/15	CLINIMIX 5/20		
Maximum Infusion Rate (mL/kg/hour)		2.4	2.4	1.67	1.25		
Corresponding	Amino Acid (g/kg/hour)	0.1*	0.1*	0.08	0.06		
infusion rate	Dextrose (g/kg/hour)	0.12	0.24	0.25*	0.25*		

<sup>\*</sup> Rate limiting factor

## 2.7 Dosage Modifications in Patients with Renal Impairment

Prior to administration, correct severe fluid or electrolyte imbalances. Closely monitor serum electrolyte levels and adjust the volume of CLINIMIX administered as required [see Warnings and Precautions (5.10)].

Patients with renal impairment not needing dialysis require 0.6 to 0.8 g of protein/kg/day. Serum electrolyte levels should be closely monitored. Patients on hemodialysis or continuous renal replacement therapy should receive 1.2 to 1.8 g of protein/kg/day up to a maximum of 2.5 g of protein/kg/day based on nutritional status and estimated protein losses. The CLINIMIX dosage can be adjusted based on the severity of renal impairment, supplementing protein as indicated. If required, additional amino acids may be added to the CLINIMIX container or infused separately. Compatibility of additions should be evaluated by a pharmacist and questions may be directed to Baxter.

# 2.8 Recommended Dosage in Pediatric Patients

The dosage and constant infusion rate of intravenous dextrose must be selected with caution in pediatric patients, particularly neonates and low weight infants, because of the increased risk of hyperglycemia/hypoglycemia [see Use in Specific Populations (8.4)]. Frequent monitoring of serum glucose concentrations is required when dextrose is prescribed to pediatric patients, particularly neonates and low birth weight infants. The infusion rate and volume should be determined by the consulting physician experienced in pediatric intravenous fluid therapy.

In pediatric patients, CLINIMIX is dosed on the basis of protein provided as amino acids. The

<sup>&</sup>lt;sup>†</sup> Protein is provided as amino acids. When infused intravenously amino acids are metabolized and utilized as the building blocks of protein.

recommended dosage, by age group is provided in **Tables 3 - 6**. Infusion rates are based on protein and do not take carbohydrates, fluid or electrolytes into consideration.

This product does not contain the amino acids cysteine and taurine, considered conditionally essential for neonates and infants. If possible, these amino acids should be added to this product if used in this pediatric population.

Table 3: Preterm and Term Infants Less than 1 Month of Age

	Recommended Nutritional	Recommended CLINIMIX Dosage in Preterm and Term Infants Less than 1 Month of Age				
	Requirements <sup>1</sup>	CLINIMIX 4.25/5	CLINIMIX4.25/10	CLINIMIX 5/15	CLINIMIX 5/20	
Infusion Rate Range (mL/kg/hr)		2.9 to 3.9	2.9 to 3.9	2.5 to 3.3	2.5 to 3.3	
Fluid (mL/kg/day)	100 to 150	70 to 94	70 to 94	60 to 79	60 to 79	
Protein* (g/kg/day) (Nitrogen g/kg/day)	3 to 4 (0.48 to 0.64)	3 to 4 (0.48 to 0.64)	3 to 4 (0.48 to 0.64)	3 to 4 (0.48 to 0.64)	3 to 4 (0.48 to 0.64)	
<b>Dextrose</b> (g/kg/day)	7 to 20	3.5 to 4.7	7 to 9.4	9 to 11.9	12 to 15.8	

<sup>\*</sup> Protein is provided as amino acids. When infused intravenously amino acids are metabolized and utilized as the building blocks of protein.

Table 4: Pediatric Patients 1 Month to Less than 1 Year of Age

	Recommended Nutritional	Recommended CLINIMIX Dosage in Pediatric Patients 1 Month to Less than 1 Year of Age				
	Requirements <sup>1</sup>	CLINIMIX 4.25/5	CLINIMIX 4.25/10	CLINIMIX 5/15	CLINIMIX 5/20	
Infusion Rate Range (mL/kg/hr)		2 to 2.9	2 to 2.9	1.7 to 2.5	1.7 to 2.5	
Fluid (mL/kg/day)	100 mL/kg for the first 10 kg + 50 mL/kg for the second 10 kg.	48 to 70	48 to 70	41 to 60	41 to 60	
Protein* (g/kg/day) (Nitrogen g/kg/day)	2 to 3 (0.32 to 0.48)	2 to 3 (0.32 to 0.48)	2 to 3 (0.32 to 0.48)	2 to 3 (0.32 to 0.48)	2 to 3 (0.32 to 0.48)	
<b>Dextrose</b> (g/kg/day)	7 to 20	2.4 to 3.5	4.8 to 7	6.1 to 9	8.2 to 12	

<sup>\*</sup> Protein is provided as amino acids. When infused intravenously amino acids are metabolized and utilized as the building blocks of protein

Table 5: Pediatric Patients 1 Year to Less than 11 Years of Age

Recommended	Recommend CLINIMIX Dosage in Pediatric Patients 1 Year to
Nutritional	Less than 11 Years of Age

	Requirements <sup>1</sup>	CLINIMIX 4.25/5	CLINIMIX 4.25/10	CLINIMIX 5/15	CLINIMIX 5/20
Infusion Rate Range (mL/kg/hr)		1 to 2	1 to 2	0.8 to 1.7	0.8 to 1.7
<b>Fluid</b> (mL/kg/day)	100 mL/kg for the first 10 kg + 50 mL/kg for the second 10 kg + 20 mL/kg for weight > 20 kg	24 to 48	24 to 48	19 to 41	19 to 41
<b>Protein</b> *(g/kg/day) (Nitrogen g/kg/day)	1 to 2 (0.16 to 0.32)	1 to 2 (0.16 to 0.32)	1 to 2 (0.16 to 0.32)	1 to 2 (0.16 to 0.32)	1 to 2 (0.16 to 0.32)
<b>Dextrose</b> (g/kg/day)	7 to 14	1.2 to 2.4	2.4 to 4.8	2.9 to 6.1	3.8 to 8.2

<sup>\*</sup> Protein is provided as amino acids. When infused intravenously amino acids are metabolized and utilized as the building blocks of protein.

Table 6: Pediatric Patients 11 Years to 17 Years of Age

	Recommended Nutritional	Recommendo	ed CLINIMIX D Years to 17	ric Patients 11	
	Requirements <sup>1</sup>	CLINIMIX 4.25/5	CLINIMIX 4.25/10	CLINIMIX 5/15	CLINIMIX 5/20
Infusion Rate Range (mL/kg/hr)		0.8 to 1.5	0.8 to 1.5	0.7 to 1.3	0.7 to 1.3
Fluid (mL/kg/day)	100 mL/kg for the first 10 kg + 50 mL/kg for the second 10 kg + 20 mL/kg for weight > 20 kg	19 to 36	19 to 36	17 to 31	17 to 31
<b>Protein</b> *(g/kg/day) (Nitrogen g/kg/day)	0.8 to 1.5 (0.13 to 0.24)	0.8 to 1.5 (0.13 to 0.24)	0.8 to 1.5 (0.13 to 0.24)	0.8 to 1.5 (0.13 to 0.24)	0.8 to 1.5 (0.13 to 0.24)
<b>Dextrose</b> (g/kg/day)	5 to 9	1 to 1.8	1.9 to 3.6	2.5 to 4.7	3.4 to 6.2

<sup>\*</sup> Protein is provided as amino acids. When infused intravenously amino acids are metabolized and utilized as the building blocks of protein.

## 2.9 Discontinuation of CLINIMIX

To reduce the risk of hypoglycemia after discontinuation, a gradual decrease in flow rate in the last hour of infusion should be considered.

## **3 DOSAGE FORMS AND STRENGTHS**

CLINIMIX is available in 1000 mL and 2000 mL dual chamber containers. The individual chambers contain essential and nonessential amino acids and dextrose. Table 7 describes the individual components of CLINIMIX.

Strength of C	LINIMIX	CLINIMIX 4.25/5 sulfiteIfree (4.25% Amino Acid in 5% Dextrose) Injection	CLINIMIX 4.25/10 sulfite Ifree (4.25% Amino Acid in 10% Dextrose) Injection	sulfite[free	CLINIMIX 5/20 sulfite free (5% Amino Acid in 20% Dextrose) Injection
	Dextrose Hydrous, USP (g/100 mL)	5	10	15	20
	Amino Acids (g/100 mL)	4.25	4.25	5	5
	Total Nitrogen (mg/100 mL)	702	702	826	826
	Leucine	311	311	365	365
	Isoleucine	255	255	300	300
	Valine	247	247	290	290
Essential Amino Acids	Lysine (added as the hydrochloride salt)	247	247	290	290
(mg/100	Phenylalanine	238	238	280	280
mL)	Histidine	204	204	240	240
	Threonine	179	179	210	210
	Methionine	170	170	200	200
	Tryptophan	77	77	90	90
	Alanine	880	880	1035	1035
Nonessentia	Arginine	489	489	575	575
Amino	Glycine	438	438	515	515
Acids	Proline	289	289	340	340
(mg/100	Serine	213	213	250	250
mL)	Tyrosine	17	17	20	20
	Acetate <sup>†</sup>	37	37	42	42
Anion Profile (mEq/L)*	Chloride <sup>‡</sup>	17	17	20	20
	pH <sup>§</sup> (Range)	6.0 (4.5 to 7.0)	6.0 (4.5 to 7.0)	6.0 (4.5 to 7.0)	6.0 (4.5 to 7.0)
	Osmolarity (mOsmol/L) (calc)	675	930	1255	1505
Caloric Content	From Dextrose	170	340	510	680
(kcal/L)	From Amino Acids	170	170	200	200
	TOTAL (Dextrose and Amino Acids)	340	510	710	880

<sup>\*</sup> Balanced by ions from amino acids.

- † Derived from glacial acetic acid (for pH adjustment) and sodium acetate.
- <sup>‡</sup> Contributed by calcium chloride, lysine hydrochloride, magnesium chloride, and sodium chloride.
- § pH of sulfite-free amino acid injection in the outlet port chamber was adjusted with glacial acetic acid.

#### **4 CONTRAINDICATIONS**

The use of CLINIMIX is contraindicated in:

- Patients with known hypersensitivity to one or more amino acids or dextrose [see Warnings and *Precautions* (5.2)].
- Patients with inborn errors of amino acid metabolism due to risk of severe metabolic and 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 parenteral nutrition. In some cases, fatal outcomes due to pulmonary embolism have occurred. CLINIMIX contains no added phosphorus. Patients, especially those with hypophosphatemia, may require the addition of phosphate. To prevent hypocalcemia, calcium supplementation should always accompany phosphate administration. Excessive addition of calcium and phosphate increases the risk of the formation of calcium phosphate precipitates. Precipitates have been reported even in the absence of phosphate salt in the solution. 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 infusion and initiate a medical evaluation. In addition to inspection of the solution [see Dosage and Administration (2.1, 2.2, 2.3, 2.4)], the infusion set and catheter should also periodically be checked for precipitates.

## 5.2 Hypersensitivity Reactions

Hypersensitivity/infusion reactions including anaphylaxis have been reported with CLINIMIX. Stop infusion immediately and treat patient accordingly if any signs or symptoms of a hypersensitivity reaction develop. Signs or symptoms may include: hypotension, hypertension, peripheral cyanosis, tachycardia, dyspnea, vomiting, nausea, urticaria, rash, pruritus, erythema, hyperhidrosis, pyrexia, and chills.

#### 5.3 Risk of Infections

Patients who require parenteral nutrition are at high risk of infections because the nutritional components of these solutions can support microbial growth. Infection and sepsis may also occur as a result of the use of intravenous catheters to administer parenteral nutrition.

The risk of infection is increased in patients with malnutrition-associated immunosuppression, hyperglycemia exacerbated by dextrose infusion, long-term use and poor maintenance of intravenous catheters, or immunosuppressive effects of other concomitant conditions, drugs, or other components of the parenteral formulation (e.g., lipid emulsion).

To decrease the risk of infection, ensure aseptic technique in catheter placement and maintenance, as well as aseptic technique in the preparation and administration of the nutritional formula.

Monitor for signs and symptoms (including fever and chills) of early infections, including laboratory test results (including leukocytosis and hyperglycemia) and frequent checks of the parenteral access device and insertion site for edema, redness and discharge.

# 5.4 Refeeding Syndrome

Refeeding severely undernourished patients may result in refeeding syndrome, characterized by the intracellular shift of potassium, phosphorus, and magnesium as the patient becomes anabolic. Thiamine deficiency and fluid retention may also develop. To prevent these complications, monitor severely undernourished patients and slowly increase nutrient intakes.

# 5.5 Hyperglycemia or Hyperosmolar Hyperglycemic State

When using CLINIMIX in patients with diabetes mellitus, impaired glucose tolerance may worsen hyperglycemia. Administration of dextrose at a rate exceeding the patient's utilization rate may lead to hyperglycemia, coma, and death. Patients with underlying confusion and renal impairment who receive dextrose infusions, may be at greater risk of developing hyperosmolar hyperglycemic state. Monitor blood glucose levels and treat hyperglycemia to maintain optimum levels while administering CLINIMIX. Insulin may be administered or adjusted to maintain optimal blood glucose levels during CLINIMIX administration.

## 5.6 Vein Damage and Thrombosis

Solutions with osmolarity of 900 mOsm/L or greater must be infused through a central catheter. CLINIMIX solutions containing more than 5% dextrose have an osmolarity greater than or equal to 900 mOsm/L. CLINIMIX 4.25/10, 5/15 and 5/20 are indicated for administration into a central vein only, such as the superior vena cava [see Dosage and Administration (2.2)]. The infusion of hypertonic nutrient injections into a peripheral vein may result in vein irritation, vein damage, and/or thrombosis.

CLINIMIX 4.25/5 is indicated for peripheral administration, or may be infused into a central vein [see Dosage and Administration (2.2)]. 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.

# 5.7 Hepatobiliary Disorders

Hepatobiliary disorders are known to develop in some patients without preexisting liver disease who receive parenteral nutrition, 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.

Increase in blood ammonia levels and hyperammonemia may occur in patients receiving amino acid solutions. In some patients this may indicate hepatic insufficiency or the presence of an inborn error of amino acid metabolism [see Contraindications (4).]

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.

#### 5.8 Aluminum Toxicity

CLINIMIX contains no more than 25 mcg/L of aluminum. However, with prolonged parenteral administration in patients with renal impairment, the aluminum contained in CLINIMIX may reach toxic levels. Preterm infants are at a greater risk because their kidneys are immature, and they require large amounts of calcium and phosphate solutions, which contain aluminum.

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

#### 5.9 Risk of Parenteral Nutrition Associated Liver Disease

Parenteral Nutrition Associated Liver Disease (PNALD) has been reported in patients who receive parenteral nutrition for extended periods of time, especially preterm infants, and can present as cholestasis or steatohepatitis. The exact etiology is unknown and is likely multifactorial. If CLINIMIX treated patients develop liver test abnormalities consider discontinuation or dosage reduction.

# 5.10 Electrolyte Imbalance and Fluid Overload

Patients with renal impairment, such as pre-renal azotemia, renal obstruction, and protein-losing nephropathy may be at increased risk of electrolyte and fluid volume imbalance. Patients with cardiac insufficiency due to left ventricular systolic dysfunction are susceptible to excess fluid accumulation. Use CLINIMIX with caution in patients with cardiac insufficiency or renal impairment. CLINIMIX dosage may require adjustment with specific attention to fluid, protein, and electrolyte content in these patients.

Monitor renal function parameters. Patients developing signs of renal impairment should be assessed early by a clinician knowledgeable in renal disease in order to determine the appropriate CLINIMIX dosage and other treatment options.

# 5.11 Monitoring/Laboratory Tests

Monitor fluid and electrolyte status, serum osmolarity, blood glucose, liver and kidney function, blood count and coagulation parameters throughout treatment.

Patients receiving CLINIMIX should be monitored frequently and their electrolyte requirements individualized.

#### **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)]
- Hypersensitivity reactions [see Warnings and Precautions (5.2)]
- Risk of Infections [see Warnings and Precautions (5.3)]
- Refeeding syndrome [see Warnings and Precautions (5.4)]
- Hyperglycemia or hyperosmolar hyperglycemic state [see Warnings and Precautions (5.5)]
- Vein damage and thrombosis [see Warnings and Precautions (5.6)]
- Hepatobiliary disorders [see Warnings and Precautions (5.7)]
- Parenteral Nutrition Associated Liver Disease [see Warnings and Precautions (5.9)]
- Electrolyte imbalance and fluid overload [see Warnings and Precautions (5.10)]

The following adverse reactions from voluntary reports or clinical studies have been reported with CLINIMIX. Because many 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.

- Diuresis
- Extravasation
- Glycosuria
- Hyperglycemia
- Hyperosmolar coma

## **8 USE IN SPECIFIC POPULATIONS**

## 8.1 Pregnancy

Risk Summary

There are no adequate or well-controlled studies in pregnant women with CLINIMIX. Additionally, animal reproduction studies have not been conducted with amino acids and electrolytes and dextrose. It

is not known whether CLINIMIX can cause fetal harm when administered to a pregnant woman.

The estimated background risk of major birth defects and miscarriage for the indicated population is unknown. However, the estimated background risk in the U.S. general population of major birth defects is 2 to 4% and of miscarriage is 15 to 20% of clinically recognized pregnancies.

### Clinical Considerations

# Disease-Associated Maternal and/or Embryo-Fetal Risk

Based on clinical practice guidelines, parenteral nutrition should be considered in cases of severe maternal malnutrition where nutritional requirements cannot be fulfilled by the enteral route because of the risks to the fetus associated with severe malnutrition, such as preterm delivery, low birth weight, intrauterine growth restriction, congenital malformations and perinatal mortality.

#### 8.2 Lactation

## Risk Summary

It is not known whether CLINIMIX is present in human milk. There are no data on the effects of CLINIMIX on the breastfed infant or on milk production. The developmental and health benefits of breastfeeding should be considered along with the mother's clinical need for CLINIMIX and any potential adverse effects on the breastfed child from CLINIMIX or from the underlying maternal condition.

#### 8.4 Pediatric Use

Safety and effectiveness of CLINIMIX in pediatric patients have not been established by adequate and well-controlled studies. Use of dextrose, amino acid infusions and electrolytes in pediatric patients is based on clinical practice [see Dosage and Administration (2.8)].

Newborns, especially those born premature and with low birth weight, are at increased risk of developing hypo – or hyperglycemia and therefore need close monitoring during treatment with intravenous glucose solutions to ensure adequate glycemic control in order to avoid potential long term adverse effects. Hypoglycemia in the newborn can cause prolonged seizures, coma and brain damage. Hyperglycemia has been associated with intraventricular hemorrhage, late onset bacterial and fungal infection, retinopathy of prematurity, necrotizing enterocolitis, bronchopulmonary dysplasia, prolonged length of hospital stay, and death. Plasma electrolyte concentrations should be closely monitored in the pediatric population as this population may have impaired ability to regulate fluids and electrolytes.

Because of immature renal function, preterm infants receiving prolonged treatment with CLINIMIX may be at risk of aluminum toxicity [see Warnings and Precautions (5.8)].

Patients, including pediatric patients, may be at risk for Parenteral Nutrition Associated Liver Disease (PNALD) [see Warnings and Precautions (5.9)].

Hyperammonemia is of special significance in infants (birth to two years). 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 (5.7)].

## 8.5 Geriatric Use

Clinical studies of CLINIMIX did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from other younger subjects. Other reported clinical experience has not identified differences in responses between the elderly and younger patients.

In general, dose selection for an elderly patient should be cautious, usually starting at the low end of the dosing range, reflecting the greater frequency of decreased hepatic, renal, or cardiac function, and of concomitant disease or drug therapy.

An increased infusion rate of CLINIMIX cause hyperglycemia, hyperosmolality, and adverse effects on water and electrolyte balance [see Warnings and Precautions (5.5, 5.10)].

Severe hyperglycemia and severe dilutional hyponatremia, and their complications, can be fatal.

Discontinue infusion and institute appropriate corrective measures in the event of overhydration or solute overload during therapy, with particular attention to respiratory and cardiovascular systems.

For current information on the management of poisoning or overdosage, contact the National Poison Control Center at 1-800-222-1222 or www.poison.org.

#### 11 DESCRIPTION

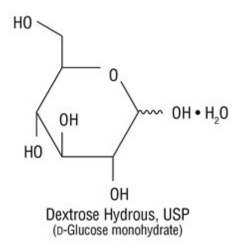
CLINIMIX sulfite-free (amino acids in dextrose) injection for intravenous use consists of sterile, nonpyrogenic, hypertonic solutions in a dual chamber container.

The outlet port chamber contains essential and nonessential amino acids. The formulas for the individual amino acids found in CLINIMIX sulfite-free (amino acids in dextrose) injections are provided in Table 8.

**Table 8: Formulas for Amino Acids** 

Essential Amino Acids	
Leucine	(CH <sub>3</sub> ) <sub>2</sub> CHCH <sub>2</sub> CH (NH <sub>2</sub> ) COOH
Isoleucine	CH <sub>3</sub> CH <sub>2</sub> CH (CH <sub>3</sub> ) CH (NH <sub>2</sub> ) COOH
Valine	(CH <sub>3</sub> ) <sub>2</sub> CHCH (NH <sub>2</sub> ) COOH
Lysine (added as the hydrochloride salt)	H <sub>2</sub> N (CH <sub>2</sub> ) <sub>4</sub> CH (NH <sub>2</sub> ) COOH
Phenylalanine	$(C_6H_5)$ CH <sub>2</sub> CH $(NH_2)$ COOH
Histadine	$(C_3H_3N_2)$ $CH_2CH$ $(NH_2)$ $COOH$
Threonine	CH <sub>3</sub> CH (OH) CH (NH <sub>2</sub> ) COO
Methionine	CH <sub>3</sub> S (CH <sub>2</sub> )2 CH (NH <sub>2</sub> ) COOH
Tryptophan	$(C_8H_6N)$ $CH_2$ $CH$ $(NH_2)$ $COOH$
Nonessential Amino Acids	
Alanine	CH <sub>3</sub> CH (NH <sub>2</sub> ) COOH
Arginine	H <sub>2</sub> NC (NH) NH (CH <sub>2</sub> )3 CH (NH <sub>2</sub> ) COOH
Glycine	H <sub>2</sub> NCH <sub>2</sub> COOH
Proline	[(CH <sub>2</sub> ) <sub>3</sub> NH CH] COOH
Serine	HOCH <sub>2</sub> CH (NH <sub>2</sub> ) COOH
Tyrosine	[C <sub>6</sub> H <sub>4</sub> (OH)] CH <sub>2</sub> CH (NH <sub>2</sub> ) COOH

The injection port chamber contains dextrose. Dextrose, USP, is chemically designated D-glucose, monohydrate ( $C_6H_{12}O_6 \cdot H_2O$ ) and has the following structure:



# 12 CLINICAL PHARMACOLOGY

#### 12.1 Mechanism of Action

CLINIMIX is used as a supplement of nutrition in patients, providing macronutrients (amino acids and dextrose) parenterally.

The amino acids provide the structural units that make up proteins and are used to synthesize proteins and other biomolecules or are oxidized to urea and carbon dioxide as a source of energy.

The administered dextrose is oxidized to carbon dioxide and water, yielding energy.

#### 12.3 Pharmacokinetics

The disposition of infused amino acids and dextrose, are essentially the same as those absorbed from ordinary food.

# **15 REFERENCES**

- 1. Ayers Phil, et al. A.S.P.E.N. Parenteral Nutrition Handbook,  $2^{nd}$  ed. 2014, pg. 123.
- 2. Mueller CM ed. *The A.S.P.E.N. Nutrition Support Core Curriculum*, 2<sup>nd</sup> ed. 2012. Chapter 29. Wolk R, Foulks C. Renal Disease, pg. 500.

## 16 HOW SUPPLIED/STORAGE AND HANDLING

CLINIMIX (amino acids in dextrose) injection (sulfite-free) is available in 1000 mL and 2000mL volumes (See Table 9).

**Table 9: CLINIMIX Formulations** 

1000 mL Code and	2000 mL Code and
NDC Number	NDC Number
Code 2B7726	Code 2B7704
NDC 033801133003	NDC 0338[]1089[]04
_	NDC Number  Code 2B7726

CLINIMIX 4.25/10 sulfite of free (4.25% Amino Acid in 10% Dextrose) Injection	Code 2B7727 NDC 0338[1134[03	Code 2B7705 NDC 0338[1091[]04
CLINIMIX 5/15 sulfitellfree (5% Amino Acid in 15% Dextrose) Injection	Code 2B7730 NDC 033801137003	Code 2B7709 NDC 0338[1099[04
CLINIMIX 5/20 sulfitellfree (5% Amino Acid in 20% Dextrose) Injection	Code 2B7731 NDC 0338[1138[]03	Code 2B7710 NDC 033801101004

Minimize exposure of CLINIMIX to heat and avoid excessive heat.

Protect from freezing.

Store CLINIMIX at room temperature (25°C/77°F) (may briefly store at up to 40°C/104°F).

Refrigerated storage is limited to 9 days once the protective clear overwrap has been opened.

Do not use if the protective clear overwrap has been previously opened or damaged.

For storage of admixed solutions see *Dosage and Administration* (2.3, 2.4).

#### 17 PATIENT COUNSELING INFORMATION

Inform patients, caregivers, or home healthcare providers of the following risks of CLINIMIX:

- Pulmonary embolism due to pulmonary vascular precipitates [see Warnings and Precautions (5.1)]
- Hypersensitivity reactions [see Warnings and Precautions (5.2)]
- Risk of Infections [see Warnings and Precautions (5.3)]
- Refeeding syndrome [see Warnings and Precautions (5.4)]
- Hyperglycemia or hyperosmolar hyperglycemic state [see Warnings and Precautions (5.5)]
- Vein damage and thrombosis [see Warnings and Precautions (5.6)]
- Hepatobiliary disorders [see Warnings and Precautions (5.7)]
- Aluminum toxicity [see Warnings and Precautions (5.8)]
- Parenteral Nutrition Associated Liver Disease (PNALD) [see Warnings and Precautions (5.9)]
- Electrolyte imbalance and fluid overload [see Warnings and Precautions (5.10)]

## **Baxter Healthcare Corporation**

Deerfield, IL 60015 USA

Printed in USA

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07-19-00-0372

### PACKAGE LABEL - PRINCIPAL DISPLAY PANEL

LOT EXP

NDC 0338-1132-03 **CLINIMIX** 2.75/5

0

0

SULFITE-FREE

(2.75% Amino Acids in 5% Dextrose) Injection

500 mL INJECTION PORT CHAMBER 10% Dextrose Injection USP

**500 mL OUTLET PORT CHAMBER** 5.5% Amino Acid Injection

2B7725

### Rx Only

0

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

	CONTENTS OF EACH 100 mL	OF THE ADMIXED	
	DEXTROSE HYDROUS USP	5 g	
	ESSENTIAL AMINO ACIDS	_	
	LEUCINE	201 mg	
	ISOLEUCINE	165 mg	
	VALINE	160 mg	
0	LYSINE (ADDED AS THE		
$\circ$	HYDROCHLORIDE SALT)	159 mg	
	PHENYLALANINE	154 mg	
	HISTIDINE	132 mg	
	THREONINE	116 mg	
	METHIONINE	110 mg	
	TRYPTOPHAN	50 mg	
	<b>HONESSENTIAL AMINO A</b>	CIDS	
	ALANINE	570 mg	
	ARGININE	316 mg	
	GLYCINE	283 mg	
	PROLINE	187 mg	
	SERINE	138 mg	
	TYROSINE	11 mg	
	mEg/L		
	ACETATE	24	
	CHLORIDE	11	
	BALANCED BY IONS FROM A	AMINO ACIDS	10
	PH ADJUSTED WITH GLACIA	L ACETIC ACID	
	STERILE		
	SINGLE DOSE CONTAINER		

0

0

Baxter

BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60016 USA MADE IN USA

#### Container Label

0

LOT EXP

2B7728 NDC 0338-1132-03

**CLINIMIX** 2.75/5 **SULFITE-FREE** 

(2.75% Amino Acid in 5% Dextrose) Injection

500 mL INJECTION PORT CHAMBER 10% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 5.5% Amino Acid Injection

## **Rx Only**

**ACTIVATE SEAL AND** MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

# A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

#### ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP 5 g

#### **ESSENTIAL AMINO ACIDS**

LEUCINE 201 mg
ISOLEUCINE 165 mg
VALINE 160 mg
LYSINE (ADDED AS THE
HYDROCHLORIDE SALT) 159 mg
PHENYLALANINE 154 mg
HISTIDINE 132 mg
THREONINE 116 mg
METHIONINE 110 mg
TRYPTOPHAN 50 mg

#### NONESSENTIAL AMINO ACIDS

ALANINE 570 mg ARGININE 316 mg GLYCINE 283 mg PROLINE 187 mg SERINE 138 mg TYROSINE 11 mg

mEq/L ACETATE 24 CHLORIDE 11

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

STERILE SINGLE DOSE CONTAINER

#### Baxter

# **BAXTER HEALTHCARE CORPORATION**

DEERFIELD IL 60015 USA MADE IN USA

LOT EXP

2B7727 NDC 0338-1134-03

# CLINIMIX 4.25/10

SULFITE-FREE

(4.25% Amino Acids in 10% Dextrose) Injection

CENTRAL LINE INFUSION ONLY 500 mL INJECTION PORT CHAMBER 20% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

Rx Only

0

0

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION DEXTROSE HYDROUS USP 10 g

ESSENTIAL AMINO ACIDS

LEUCINE 311 mg

LEUCINE 311 mg
ISOLEUCINE 255 mg
VALINE 247 mg

LYSINE (ADDED AS THE
HYDRÔCHLORIDE SALT) 247 mg

0

0

0

| 17 mg | 17 m

NONESSENTIAL AMINO ACIDS

ALANINE 890 mg
ARGININE 499 mg
GLYCINE 438 mg
PROLINE 289 mg
SERINE 213 mg
TYROSINE 17 mg

mEq/L ACETATE 37 CHLORIDE 17

BALANCED BY IONS FROM AMINO ACIDS ph adjusted with Glacial acetic acid sterile

SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F) AVOID EXCESSIVE HEAT PROTECT FROM FREEZING SEE PRESCRIBING INFORMATION

BAXTER BAXTER HEALTHCARE CORPORATION DEERFELD IL 60015 USA MADE N USA

O O Container Label

0

LOT EXP 2B7727 NDC 0338-1134-03

CLINIMIX 4.25/10 SULFITE-FREE

(4.25% Amino Acids in 10% Dextrose) Injection

500 mL INJECTION PORT CHAMBER 20% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

Rx Only

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP 10 g

## **ESSENTIAL AMINO ACIDS**

LEUCINE 311 mg

ISOLEUCINE 255 mg

VALINE 247 mg

LYSINE (ADDED AS THE

HYDROCHLORIDE SALT) 247 mg

PHENYLALANINE 238 mg

HISTIDINE 204 mg

THREONINE 179 mg

METHIONINE 170 mg

TRYPTOPHAN 77 mg

### NONESSENTIAL AMINO ACIDS

ALANINE 880 mg

ARGININE 489 mg

GLYCINE 438 mg

PROLINE 289 mg

SERINE 213 mg

TYROSINE 17 mg

mEq/L

**ACETATE 37** 

**CHLORIDE 17** 

BALANCED BY IONS FROM AMINO ACIDS

pH ADJUSTED WITH GLACIAL ACETIC ACID

**STERILE** 

SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F) AVOID EXCESSIVE HEAT PROTECT FROM FREEZING SEE PRESCRIBING INFORMATION

Baxter

## **BAXTER HEALTHCARE CORPORATION**

DEERFIELD IL 60015 USA

MADE IN USA

				LOT	EXP		
	2B7701 NDC 0338-1083-04			CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION			
				DEXTROSE HYDROUS USP	5 g		
)	2.75/5 CLINIMIX 2.75/5 sulfite-free (2.75% Amino Acid	0	0	ESSENTIAL AMINO ACIDS LEUCINE ISOLEUCINE VALINE LYSINE (ADDED AS THE HYDROCHLORIDE SALT) PHENYLALANINE HISTIDINE THREONINE	201 mg 165 mg 160 mg 159 mg 154 mg 132 mg 116 mg	_	0
	in 5% Dextrose)			METHIONINE TRYPTOPHAN	110 mg 50 mg		
	Injection  1000 mL INJECTION PORT CHAMBER 10% Dextrose Injection USP			NONESSENTIAL AMINO ACI ALANINE ARGININE GLYCINE PROLINE SERINE			
	10 /6 Dextrose Injection our			TYROSINE	11 mg	10 (0)	
	1000 mL OUTLET PORT CHAMBER 5.5% Amino Acid Injection			<b>mEq/L</b> ACETATE 24 CHLORIDE 11		er en	
	200200	0	0	BALANCED BY IONS FROM AMI	INO ACIDS		0
	Rx Only	0	0	PH ADJUSTED WITH GLACIAL A			0
	CHECK FOR MINUTE LEAKS BY SQUEEZING EACH CHAMBER OF THE BAG			PH 6.0 (4.5 TO 7.0) HYPERTON OSMOLARITY 525 mOsmol/L (C.			
	BEFORE USE GRASP EACH SIDE OF THE TOP OF THE BAG AND ROLL BAG TO OPEN SEAL BETWEEN CHAMBERS			STERILE NONPYROGENIC SINGLE DOSE CONTAINER			
	MIX THOROUGHLY						
	AFTER MIXING THE PRODUCT REPRESENTS 2000 mL					W - W	
	ADDITIVES MAY BE INCOMPATIBLE CONSULT WITH PHARMACIST IF AVAILABLE WHEN INTRODUCING ADDITIVES USE ASEPTIC TECHNIQUE MIX THOROUGHLY DO NOT STORE						
	DOSAGE INTRAVENOUSLY AS DIRECTED BY A PHYSICIAN DO NOT USE UNLESS SOLUTION IS CLEAR REFRIGERATED STORAGE AFTER MIXING LIMITED TO 24 HOURS DISCARD UNUSED PORTION SEE ACCOMPANYING DIRECTIONS FOR USE	0	0	Baxter			0
	CAUTIONS MUST NOT BE USED IN SERIES CONNECTIONS DISCONTINUE INFUSION IF ADVERSE REACTION OCCURS	_	_	BAXTER HEALTHCARE CORPORATION CLINTEC NUTRITION DIVISION DEERFIELD IL 80015 USA MADE IN USA PATENT PENDING	N		
	CLARITY DUAL CHAMBER CONTAINER PL 2401 PLASTIC			BAXTER CLINIMIX AND CLARITY ARE TO OF BAXTER INTERNATIONAL INC	RADEMARKS		
		Containe	r L	abel			

#### **Container Label**

## LOT EXP

O Rx Only

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2B7701 NDC 0338-1083-04

**CLINIMIX 2.75/5** sulfite-free (2.75% Amino Acid in 5% Dextrose) Injection

1000 mL INJECTION PORT CHAMBER 10% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 5.5% Amino Acid Injection

# **Rx Only**

CHECK FOR MINUTE LEAKS BY SQUEEZING EACH CHAMBER OF THE BAG BEFORE USE GRASP EACH SIDE OF

THE TOP OF THE BAG AND ROLL BAG

TO OPEN SEAL BETWEEN CHAMBERS

MIX THOROUGHLY

AFTER MIXING THE PRODUCT

REPRESENTS 2000 mL

ADDITIVES MAY BE INCOMPATIBLE CONSULT WITH

PHARMACIST IF AVAILABLE WHEN INTRODUCING

ADDITIVES USE ASEPTIC TECHNIQUE

MIX THOROUGHLY DO NOT STORE

DOSAGE INTRAVENOUSLY AS DIRECTED BY A

PHYSICIAN DO NOT USE UNLESS SOLUTION IS

CLEAR REFRIGERATED STORAGE AFTER MIXING

LIMITED TO 24 HOURS DISCARD UNUSED PORTION

SEE ACCOMPANYING DIRECTIONS FOR USE

CAUTIONS MUST NOT BE USED IN SERIES

CONNECTIONS DISCONTINUE INFUSION

IF ADVERSE REACTION OCCURS

CLARITY DUAL CHAMBER CONTAINER PL 2401 PLASTIC

CONTENTS OF EACH 100 mL OF THE

ADMIXED INJECTION

DEXTROSE HYDROUS USP 5 g

#### **ESSENTIAL AMINO ACIDS**

LEUCINE 201 mg

**ISOLEUCINE 165 mg** 

VALINE 160 mg

LYSINE (ADDED AS THE

HYDROCHLORIDE SALT) 159 mg

PHENYLALANINE 154 mg

HISTIDINE 132 mg

THREONINE 116 mg

METHIONINE 110 mg

TRYPTOPHAN 50 mg

## NONESSENTIAL AMINO ACIDS

ALANINE 570 mg

ARGININE 316 mg

GLYCINE 283 mg

PROLINE 187 mg

SERINE 138 mg

TYROSINE 11 mg

mEq/L

**ACETATE 24** 

**CHLORIDE 11** 

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

pH 6.0 (4.5 TO 7.0) HYPERTONIC OSMOLARITY 525 mOsmol/L (CALC) STERILE NONPYROGENIC SINGLE DOSE CONTAINER

Baxter

0

BAXTER HEALTHCARE CORPORATION
CLINTEC NUTRITION DIVISION
DEERFIELD IL 60015 USA
MADE IN USA
PATENT PENDING
BAXTER CLINIMIX AND CLARITY ARE TRADEMARKS
OF BAXTER INTERNATIONAL INC

2B7726 NDC 0338-1133-03

4.25/5

SULFITE-FREE

(4.25% Amino Acids in 5% Dextrose) Injection

0

0

500 mL INJECTION PORT CHAMBER 10% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

O Rx Only

0

4.25/5

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

LOT EXP

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION DEXTROSE HYDROUS USP 5 g

ESSENTIAL AMINO ACIDS

 LEUCINE
 311 mg

 ISOLEUCINE
 255 mg

 VALINE
 247 mg

0

0

0

LYSINE (ADDED AS THE
HYDRÓCHLORIDE SALT) 247 mg
PHENYLALANINE 238 mg
HISTIDINE 204 mg
THREONINE 179 mg

 THREONINE
 179 mg

 METHIONINE
 170 mg

 TRYPTOPHAN
 77 mg

 NONESSENTIAL AMINO ACIDS

 ALANINE
 880 mg

 ARGININE
 489 mg

 GLYCINE
 438 mg

 PROLINE
 300 mg

 PROLINE
 289 mg

 SERINE
 213 mg

 TYROSINE
 17 mg

mEq/L
ACETATE 37
CHLORIDE 17

BALANCED BY IONS FROM AMINO ACIDS PH ADJUSTED WITH GLACIAL ACETIC ACID

STERILE SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F) AVOID EXCESSIVE HEAT

PROTECT FROM FREEZING SEE PRESCRIBING INFORMATION

BAXTER HEALTHCARE CORPORATION DEERFELD IL 60016 USA

Container Label

LOT EXP 2B7726 NDC 0338-1133-03 CLINIMIX **SULFITE-FREE** 

(4.25% Amino Acids in 5% Dextrose)
Injection

500 mL INJECTION PORT CHAMBER 10% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

**ACTIVATION** 

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED

**INJECTION** 

DEXTROSE HYDROUS USP 5 g

#### **ESSENTIAL AMINO ACIDS**

LEUCINE 311 mg

ISOLEUCINE 255 mg

VALINE 247 mg

LYSINE (ADDED AS THE

HYDROCHLORIDE SALT) 247 mg

PHENYLALANINE 238 mg

HISTIDINE 204 mg

THREONINE 179 mg

METHIONINE 170 mg

TRYPTOPHAN 77 mg

#### NONESSENTIAL AMINO ACIDS

ALANINE 880 mg

ARGININE 489 mg

GLYCINE 438 mg

PROLINE 289 mg

SERINE 213 mg

TYROSINE 17 mg

mEq/L

**ACETATE 37** 

**CHLORIDE 17** 

BALANCED BY IONS FROM AMINO ACIDS

pH ADJUSTED WITH GLACIAL ACETIC ACID

**STERILE** 

SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F) AVOID EXCESSIVE HEAT PROTECT FROM FREEZING SEE PRESCRIBING INFORMATION

#### Baxter

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BAXTER HEALTHCARE CORPORATION **DEERFIELD IL 60015 USA** MADE IN USA

> 2B7704 NDC 0338-1089-04 CLINIMIX 4.25/5

> > SULFITE-FREE

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(4.25% Amino Acids in 5% Dextrose) Injection

1000 mL INJECTION PORT CHAMBER 10% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

Rx Only

**ACTIVATE SEAL AND** MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

LOT EXP

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP

311 mg

0

0

0

ESSENTIAL AMINO ACIDS

ISOLEUCINE 255 mg VALINE 247 mg LYSINE (ADDED AS THE HYDROCHLORIDE SALT) 247 mg 238 mg PHENYLALANINE 204 mg HISTIDINE THREONINE 179 mg METHIONINE

NONESSENTIAL AMINO ACIDS

TRYPTOPHAN

ALANINE 880 mg ARGININE 489 mg GLYCINE 438 mg PROLINE 289 mg SERINE 213 mg TYROSINE 17 mg

mEq/L ACETATE 37 CHLORIDE 17

BALANCED BY IONS FROM AMINO ACIDS PH ADJUSTED WITH GLACIAL ACETIC ACID

STERILE SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F)

AVOID EXCESSIVE HEAT

PROTECT FROM FREEZING

SEE PRESCRIBING INFORMATION

Baxter BAXTER HEALTHCARE CORPORATION DEERFELD L 60016 USA

**Container Label** 

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#### 2B7704 NDC 0338-1089-04

CLINIMIX 4.25/5 SULFITE-FREE (4.25% Amino Acid in 5% Dextrose) Injection

1000 mL INJECTION PORT CHAMBER 10% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

## **Rx Only**

# ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP 5 g

#### **ESSENTIAL AMINO ACIDS**

LEUCINE 311 mg
ISOLEUCINE 255 mg
VALINE 247 mg
LYSINE (ADDED AS THE
HYDROCHLORIDE SALT) 247 mg
PHENYLALANINE 238 mg
HISTIDINE 204 mg
THREONINE 179 mg
METHIONINE 170 mg
TRYPTOPHAN 77 mg

## NONESSENTIAL AMINO ACIDS

ALANINE 880 mg ARGININE 489 mg GLYCINE 438 mg PROLINE 289 mg SERINE 213 mg TYROSINE 17 mg

mEq/L ACETATE 37 CHLORIDE 17

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

STERILE SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F) AVOID EXCESSIVE HEAT PROTECT FROM FREEZING SEE PRESCRIBING INFORMATION

#### Baxter

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BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA

2B7705 NDC 0338-1091-04

# CLINIMIX 4.25/10

SULFITE-FREE

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(4.25% Amino Acids in 10% Dextrose) Injection

## CENTRAL LINE INFUSION ONLY

1000 mL INJECTION PORT CHAMBER 20% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

**Rx Only** 

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

LOT

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

EXP

311 mg

247 mg

0

0

0

DEXTROSE HYDROUS USP

ESSENTIAL AMINO ACIDS

LEUCINE

ISOLEUCINE 255 mg VALINE 247 mg

LYSINE (ADDED AS THE HYDROCHLORIDE SALT)

PHENYLALANINE 238 mg HISTIDINE 204 mg

 THREONINE
 179 mg

 METHIONINE
 170 mg

 TRYPTOPHAN
 77 mg

NONESSENTIAL AMINO ACIDS

ALANINE 880 mg
ARGININE 489 mg
GLYCINE 438 mg
PROLINE 289 mg

SERINE 213 mg
TYROSINE 17 mg

mEq/L ACETATE 37

CHLORIDE 17

BALANCED BY IONS FROM AMINO ACIDS

PH ADJUSTED WITH GLACIAL ACETIC ACID STERILE

SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F)

AVOID EXCESSIVE HEAT

PROTECT FROM FREEZING

SEE PRESCRIBING INFORMATION

Baxter BAXTER HEALTHCAR

BAXTER HEALTHCARE CORPORATION DEERFELD L 60016 USA

Container Label

LOT EXP

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2B7705 NDC 0338-1091-04

CLINIMIX 4.25/10 SULFITE-FREE (4.25% Amino Acid

# in 10% Dextrose) Injection

1000 mL INJECTION PORT CHAMBER 20% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

## Rx Only

## ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

**ACTIVATION** 

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP 10 g

## **ESSENTIAL AMINO ACIDS**

LEUCINE 311 mg

ISOLEUCINE 255 mg

VALINE 247 mg

LYSINE (ADDED AS THE

HYDROCHLORIDE SALT) 247 mg

PHENYLALANINE 238 mg

HISTIDINE 204 mg

THREONINE 179 mg

METHIONINE 170 mg

TRYPTOPHAN 77 mg

# **NONESSENTIAL AMINO ACIDS**

ALANINE 880 mg

ARGININE 489 mg

GLYCINE 438 mg

PROLINE 289 mg

SERINE 213 mg

TYROSINE 17 mg

mEq/L

ACETATE 37

**CHLORIDE 17** 

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

STERILE

SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F)

AVOID EXCESSIVE HEAT

# PROTECT FROM FREEZING SEE PRESCRIBING INFORMATION

#### Baxter

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BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA

2B7728

NDC 0338-1135-03

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# CLINIMIX 4.25/20

SULFITE-FREE

(4.25% Amino Acids in 20% Dextrose) Injection

## **CENTRAL LINE INFUSION ONLY**

500 mL INJECTION PORT CHAMBER 40% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

Rx Only

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

LOT

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION
DEXTROSE HYDROUS USP 20 g

EXP

0

0

ESSENTIAL AMINO ACIDS

LEUCINE 311 mg
isoLeucine 255 mg
VALINE 247 mg
LYSINE (ADDED AS THE
HYDROCHLORIDE SALT) 247 mg

 PHENYLALANINE
 238 mg

 HISTIDINE
 204 mg

 THREONINE
 179 mg

 METHIONINE
 170 mg

 TRYPTOPHAN
 77 mg

NONESSENTIAL AMINO ACIDS

ALANINE 880 mg
ARGININE 489 mg
GLYCINE 438 mg
PROLINE 289 mg
SERINE 213 mg
TYROSINE 17 mg

mEq/L
ACETATE 37
CHLORIDE 17

BALANCED BY IONS FROM AMINO ACIDS ph adjusted with Glacial acetic acid sterile

→ SINGLE DOSE CONTAINER

Baxter
BAXTER HEALTHCARE CORPORATION
DEERFIELD IL 60016 USA

**Container Label** 

#### LOT EXP

2B7728 NDC 0338-1035-03

CLINIMIX
4.25/20
SULFITE-FREE
(4.25% Amino Acid
in 20% Dextrose)

500 mL INJECTION PORT CHAMBER 40% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

**Rx Only** 

Injection

# ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

**ACTIVATION** 

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP 20 g

#### **ESSENTIAL AMINO ACIDS**

LEUCINE 311 mg
ISOLEUCINE 255 mg
VALINE 247 mg
LYSINE (ADDED AS THE
HYDROCHLORIDE SALT) 247 mg
PHENYLALANINE 238 mg
HISTIDINE 204 mg
THREONINE 179 mg
METHIONINE 170 mg

### NONESSENTIAL AMINO ACIDS

ALANINE 880 mg ARGININE 489 mg GLYCINE 438 mg PROLINE 289 mg SERINE 213 mg TYROSINE 17 mg

TRYPTOPHAN 77 mg

mEq/L

ACETATE 37 CHLORIDE 17

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

STERILE

SINGLE DOSE CONTAINER

#### Baxter

BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA LOT EXP

CLINIMIX 4.25/20

NDC 0338-1093-04

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SULFITE-FREE

(4.25% Amino Acids in 20% Dextrose) Injection

### CENTRAL LINE INFUSION ONLY

2B7706

1000 mL INJECTION PORT CHAMBER 40% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

Rx Only

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ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP 20 a

ESSENTIAL AMINO ACIDS

LEUCINE 311 mg ISOLEUCINE 255 mg

0

0

VALINE 247 mg
LYSINE (ADDED AS THE
HYDROCHLORIDE SALT) 247 mg
PHENYLALANINE 238 mg

 PHENYLALANINE
 238 mg

 HISTIDINE
 204 mg

 THREONINE
 179 mg

 METHONINE
 170 mg

 TRYPTOPHAN
 77 mg

NONESSENTIAL AMINO ACIDS

ALANINE 880 mg
ARGININE 489 mg
GLYCINE 488 mg
PROLINE 289 mg
SERINE 213 mg
TYROSINE 17 mg

mEq/L ACETATE 37

BALANCED BY IONS FROM AMINO ACIDS ph adjusted with Glacial Acetic acid

STERILE SINGLE DOSE CONTAINER

CHLORIDE

Baxter

BAXTER HEALTHCARE CORPORATION DEEPS ELD IL 60015 USA WADE IN USA

#### **Container Label**

#### LOT EXP

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2B7706 NDC 0338-1093-04

CLINIMIX 4.25/20 SULFITE-FREE (4.25% Amino Acid in 20% Dextrose) Injection

1000 mL INJECTION PORT CHAMBER 40% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

#### Rx Only

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE

**ADMIXED INJECTION** 

DEXTROSE HYDROUS USP 20 g

## **ESSENTIAL AMINO ACIDS**

LEUCINE 311 mg

ISOLEUCINE 255 mg

VALINE 247 mg

LYSINE (ADDED AS THE

HYDROCHLORIDE SALT) 247 mg

PHENYLALANINE 238 mg

HISTIDINE 204 mg

THREONINE 179 mg

METHIONINE 170 mg

TRYPTOPHAN 77 mg

## NONESSENTIAL AMINO ACIDS

ALANINE 880 mg

ARGININE 489 mg

GLYCINE 438 mg

PROLINE 289 mg

SERINE 213 mg

TYROSINE 17 mg

mEq/L

**ACETATE 37** 

**CHLORIDE 17** 

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

**STERILE** 

SINGLE DOSE CONTAINER

Baxter

BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA

LOT EXP

2B7729 NDC 0338-1136-03 CLINIMIX 4.25/25

SULFITE-FREE

(4.25% Amino Acids in 25% Dextrose)
Injection

# CENTRAL LINE INFUSION ONLY

500 mL INJECTION PORT CHAMBER 50% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

O Rx Only

0

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

	CONTENTS OF EACH 100 mL INJECTION DEXTROSE HYDROUS USP	OF THE ADMIXED		
	ESSENTIAL AMINO ACIDS			
	LEUCINE	311 mg		
	ISOLEUCINE	255 mg		
	VALINE	247 mg		
)	LYSINE (ADDED AS THE			$\circ$
	HYDROCHLORIDE SALT)	247 mg		_
	PHENYLALANINE	238 mg		
	HISTIDINE	204 mg		
	THREONINE	179 mg		
	METHIONINE	170 mg		
	TRYPTOPHAN	77 mg		
	NONESSENTIAL AMINO A	CIDS		
	ALANINE	980 mg		
	ARGININE	489 mg		
	GLYCINE	438 mg		
	PROLINE	289 mg		
	SERINE	213 mg		
	TYROSINE	17 mg		
	mEq/L			
	ACETATE	37		
	CHLORIDE	17		
	BALANCED BY IONS FROM AMINO ACIDS			
	PH ADJUSTED WITH GLACIAL ACETIC ACID			
	STERILE			
١	SINGLE DOSE CONTAINER			0

Baxter BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA

# **Container Label**

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## LOT EXP

2B7729 NDC 0338-1136-03

CLINIMIX 4.25/25 SULFITE-FREE (4.25% Amino Acid in 25% Dextrose) Injection

500 mL INJECTION PORT CHAMBER 50% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

#### Rx Only

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

**ACTIVATION** 

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

## A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE

ADMIXED INJECTION

DEXTROSE HYDROUS USP 25 g

#### **ESSENTIAL AMINO ACIDS**

LEUCINE 311 mg

ISOLEUCINE 255 mg

VALINE 247 mg

LYSINE (ADDED AS THE

HYDROCHLORIDE SALT) 247 mg

PHENYLALANINE 238 mg

HISTIDINE 204 mg

THREONINE 179 mg

METHIONINE 170 mg

TRYPTOPHAN 77 mg

## NONESSENTIAL AMINO ACIDS

ALANINE 880 mg

ARGININE 489 mg

GLYCINE 438 mg

PROLINE 289 mg

SERINE 213 mg

TYROSINE 17 mg

mEq/L

**ACETATE 37** 

**CHLORIDE 17** 

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

**STERILE** 

SINGLE DOSE CONTAINER

#### Baxter

BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA LOT EXP

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

2B7707 NDC 0338-1095-04

**CLINIMIX** 

4.25/25

0

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0

SULFITE-FREE

(4.25% Amino Acids in 25% Dextrose) Injection

#### CENTRAL LINE INFUSION ONLY

1000 mL INJECTION PORT CHAMBER 50% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER

○ 8.5% Amino Acid Injection

**Rx Only** 

0

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

DEXTROSE HYDROUS USP 25 g **ESSENTIAL AMINO ACIDS** LEUCINE 311 mg ISOLEUCINE VALINE LYSINE (ADDED AS THE HYDROCHLORIDE SALT) 247 mg PHENYLALANINE 239 mg 204 mg HISTIDINE THREONINE 179 mg METHIONINE 170 mg TRYPTOPHAN NONESSENTIAL AMINO ACIDS

0

0

0

mEq/L ACETATE 37 CHLORIDE 17

BALANCED BY IONS FROM AMINO ACIDS

ph ADJUSTED WITH GLACIAL ACETIC ACID

STERILE
SINGLE DOSE CONTAINER

BAXTER
BAXTER HEALTHCARE CORPORATION
DEFFELD IL 80018 USA
MADE IN USA

#### **Container Label**

LOT EXP

0

2B7707 NDC 0338-1095-04

CLINIMIX
4.25/25
SULFITE-FREE
(4.25% Amino Acid
in 25% Dextrose)
Injection

1000 mL INJECTION PORT CHAMBER 50% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 8.5% Amino Acid Injection

Rx Only

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE

ADMIXED INJECTION

DEXTROSE HYDROUS USP 25 g

### **ESSENTIAL AMINO ACIDS**

LEUCINE 311 mg

ISOLEUCINE 255 mg

VALINE 247 mg

LYSINE (ADDED AS THE

HYDROCHLORIDE SALT) 247 mg

PHENYLALANINE 238 mg

HISTIDINE 204 mg

THREONINE 179 mg

METHIONINE 170 mg

TRYPTOPHAN 77 mg

### NONESSENTIAL AMINO ACIDS

ALANINE 880 mg

ARGININE 489 mg

GLYCINE 438 mg

PROLINE 289 mg

SERINE 213 mg

TYROSINE 17 mg

mEq/L

**ACETATE 37** 

**CHLORIDE 17** 

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

**STERILE** 

SINGLE DOSE CONTAINER

**Baxter** 

BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA 2B7730 NDC 0338-1137-03

5/15

SULFITE-FREE

0

0

0

(5% Amino Acids in 15% Dextrose) Injection

### CENTRAL LINE INFUSION ONLY

500 mL INJECTION PORT CHAMBER 30% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 10% Amino Acid Injection

→ Rx Only

0

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF FACILIES -1 OF THE ADMINED

LOT

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP 15 a

EXP

90 mg

0

0

0

ESSENTIAL AMINO ACIDS
LEUCINE 365 mg
ISOLEUCINE 300 mg

VALINE 290 mg
LYSINE (ADDED AS THE
HYDROCHLORIDE SALT) 290 mg
PHENYLALANINE 280 mg
HISTIDINE 240 mg
THREONINE 210 mg
METHIONINE 200 mg

NONESSENTIAL AMINO ACIDS

TRYPTOPHAN

ALANINE 1035 mg
ARGININE 575 mg
GLYCINE 515 mg
PROLINE 340 mg
SERINE 250 mg
TYROSINE 20 mg

mEq/L ACETATE 42 CHLORIDE 20

BALANCED BY IONS FROM AMINO ACIDS PH ADJUSTED WITH GLACIAL ACETIC ACID STERILE

SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F) AVOID EXCESSIVE HEAT PROTECT FROM FREEZING SEE PRESCRIBING INFORMATION

Baxter

BAXTER HEALTHCARE CORPORATION DEERFELD IL 60016 USA

O Container Label

LOT EXP

0

2B7730 NDC 0338-1137-03

CLINIMIX 5/15 SULFITE-FREE (5% Amino Acid in 15% Dextrose)

Injection

500 mL INJECTION PORT CHAMBER 30% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 10% Amino Acid Injection

Rx Only

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL

### REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE

**ADMIXED INJECTION** 

DEXTROSE HYDROUS USP 15 g

#### **ESSENTIAL AMINO ACIDS**

LEUCINE 365 mg

ISOLEUCINE 300 mg

VALINE 290 mg

LYSINE (ADDED AS THE

HYDROCHLORIDE SALT) 290 mg

PHENYLALANINE 280 mg

HISTIDINE 240 mg

THREONINE 210 mg

METHIONINE 200 mg

TRYPTOPHAN 90 mg

### NONESSENTIAL AMINO ACIDS

ALANINE 1035 mg

ARGININE 575 mg

GLYCINE 515 mg

PROLINE 340 mg

SERINE 250 mg

TYROSINE 20 mg

mEq/L

**ACETATE 42** 

**CHLORIDE 20** 

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

**STERILE** 

SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F)

AVOID EXCESSIVE HEAT

PROTECT FROM FREEZING

SEE PRESCRIBING INFORMATION

**Baxter** 

BAXTER HEALTHCARE CORPORATION

**DEERFIELD IL 60015 USA** 

MADE IN USA

LOT EXP

2B7709 NDC 0338-1099-04 CLINIMIX 5/15

#### SULFITE-FREE

0

0

0

(5% Amino Acids in 15% Dextrose) Injection

### CENTRAL LINE INFUSION ONLY

1000 mL INJECTION PORT CHAMBER 30% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 10% Amino Acid Injection

### Rx Only

0

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON AFTER MIXING THE PRODUCT REPRESENTS 2000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION DEXTROSE HYDROUS USP

365 mg

0

0

0

**ESSENTIAL AMINO ACIDS** LEUCINE

ISOLEUCINE 300 mg VALINE 290 mg LYSINE (ADDED AS THE HYDROCHLORIDE SALT) 290 mg PHENYLALANINE 280 mg HISTIDINE 240 mg THREONINE 210 mg METHIONINE 200 mg TRYPTOPHAN 90 mg

NONESSENTIAL AMINO ACIDS

ALANINE 1035 mg ARGININE 575 mg GLYCINE 515 mg 340 mg PROUNE 250 mg SERINE TYROSINE 20 mg

mEq/L **ACETATE** 

42 CHLORIDE 20

BALANCED BY IONS FROM AMINO ACIDS

PH ADJUSTED WITH GLACIAL ACETIC ACID

STERILE SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F) AVOID EXCESSIVE HEAT

PROTECT FROM FREEZING

SEE PRESCRIBING INFORMATION

Baxter BAXTER HEALTHCARE CORPORATION DEERFELD IL 60016 USA

#### **Container Label**

LOT EXP

0

2B7709 NDC 0338-1099-04

**CLINIMIX** 5/15 SULFITE-FREE (5% Amino Acid in 15% Dextrose) Injection

1000 mL INJECTION PORT CHAMBER 30% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 10% Amino Acid Injection

Rx Only

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

#### **ACTIVATION**

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP 15 g

#### **ESSENTIAL AMINO ACIDS**

LEUCINE 365 mg
ISOLEUCINE 300 mg
VALINE 290 mg
LYSINE (ADDED AS THE
HYDROCHLORIDE SALT) 290 mg
PHENYLALANINE 280 mg
HISTIDINE 240 mg
THREONINE 210 mg
METHIONINE 200 mg

### **NONESSENTIAL AMINO ACIDS**

ALANINE 1035 mg ARGININE 575 mg GLYCINE 515 mg PROLINE 340 mg SERINE 250 mg TYROSINE 20 mg

TRYPTOPHAN 90 mg

mEq/L

ACETATE 42 CHLORIDE 20

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

**STERILE** 

SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F) AVOID EXCESSIVE HEAT PROTECT FROM FREEZING SEE PRESCRIBING INFORMATION

### Baxter

BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA

LOT EXP

2B7731 NDC 0338-1138-03 **CLINIMIX** 5/20

SULFITE-FREE

0

0

0

(5% Amino Acids in 20% Dextrose) Injection

### CENTRAL LINE INFUSION ONLY

500 mL INJECTION PORT CHAMBER 40% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 10% Amino Acid Injection

O Rx Only

0

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED DEXTROSE HYDROUS USP

365 mg

0

0

0

ESSENTIAL AMINO ACIDS LEUCINE

ISOLEUCINE 300 mg VALINE LYSINE (ADDED AS THE HYDROCHLORIDE SALT) 290 mg 290 mg PHENYLALANINE 280 mg HISTIDINE 240 mg THREONINE 210 mg METHIONINE 200 mg 90 mg TRYPTOPHAN

NONESSENTIAL AMINO ACIDS

1035 mg 575 mg ALANINE ARGININE GLYCINE 515 mg PROLINE 340 mg 250 mg 20 mg TYROSINE

mEq/L ACETATE

CHLORIDE BALANCED BY IONS FROM AMINO ACIDS

PH ADJUSTED WITH GLACIAL ACETIC ACID STERILE

SINGLE DOSE CONTAINER ROOM TEMPERATURE (25°C/77°F)

AVOID EXCESSIVE HEAT

PROTECT FROM FREEZING

SEE PRESCRIBING INFORMATION

Baxter

BAXTER HEALTHCARE CORPORATION DEERFELD IL 60016 USA

**Container Label** 

0

## **LOT EXP**

2B7731 NDC 0338-1138-03

**CLINIMIX** 

5/

0

**SULFITE-FREE** 

(5% Amino Acid

in 20% Dextrose)

Injection

500 mL INJECTION PORT CHAMBER 40% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER

10% Amino Acid Injection

Rx Only

**ACTIVATE SEAL AND** MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON **ACTIVATION** 

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE

**ADMIXED INJECTION** 

DEXTROSE HYDROUS USP 20 g

#### **ESSENTIAL AMINO ACIDS**

LEUCINE 365 mg

ISOLEUCINE 300 mg

VALINE 290 mg

LYSINE (ADDED AS THE

HYDROCHLORIDE SALT) 290 mg

PHENYLALANINE 280 mg

HISTIDINE 240 mg

THREONINE 210 mg

METHIONINE 200 mg

TRYPTOPHAN 90 mg

### NONESSENTIAL AMINO ACIDS

ALANINE 1035 mg

ARGININE 575 mg

GLYCINE 515 mg

PROLINE 340 mg

SERINE 250 mg

TYROSINE 20 mg

mEq/L

**ACETATE 42** 

**CHLORIDE 20** 

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

**STERILE** 

SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F)

AVOID EXCESSIVE HEAT

PROTECT FROM FREEZING

SEE PRESCRIBING INFORMATION

#### Baxtei

BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA

LOT EXP

2B7710 NDC 0338-1101-04
CLINIMIX
5/20

#### SULFITE-FREE

0

0

0

(5% Amino Acids in 20% Dextrose) Injection

### CENTRAL LINE INFUSION ONLY

1000 mL INJECTION PORT CHAMBER 40% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 10% Amino Acid Injection

### Rx Only

0

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP 20 g

ESSENTIAL AMINO ACIDS

 LEUCINE
 365 mg

 ISOLEUCINE
 300 mg

 VALINE
 290 mg

LYSINE (ADDED AS THE
HYDROCHLORIDE SALT) 290 mg
PHENYLALANINE 280 mg
HISTIDINE 240 mg
THREONINE 210 mg
METHIONINE 200 mg

NONESSENTIAL AMINO ACIDS

TRYPTOPHAN

ALANINE 1035 mg
ARGININE 575 mg
GLYCINE 515 mg
PROLINE 340 mg
SERINE 250 mg
TYROSINE 20 mg

90 mg

0

0

mEq/L ACETATE 42 CHLORIDE 20

BALANCED BY IONS FROM AMINO ACIDS

PH ADJUSTED WITH GLACIAL ACETIC ACID STERILE

SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F) AVOID EXCESSIVE HEAT PROTECT FROM FREEZING SEE PRESCRIBING INFORMATION

BAXTER HEALTHCARE CORPORATION DEERFELD IL 60016 USA MADE N USA

#### **Container Label**

LOT EXP

0

2B7710 NDC 0338-1101-04

CLINIMIX 5/20 SULFITE-FREE (5% Amino Acid in 20% Dextrose) Injection

1000 mL INJECTION PORT CHAMBER 40% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 10% Amino Acid Injection

Rx Only

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

#### **ACTIVATION**

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP 20 g

#### **ESSENTIAL AMINO ACIDS**

LEUCINE 365 mg
ISOLEUCINE 300 mg
VALINE 290 mg
LYSINE (ADDED AS THE
HYDROCHLORIDE SALT) 290 mg
PHENYLALANINE 280 mg
HISTIDINE 240 mg
THREONINE 210 mg
METHIONINE 200 mg

### **NONESSENTIAL AMINO ACIDS**

ALANINE 1035 mg ARGININE 575 mg GLYCINE 515 mg PROLINE 340 mg SERINE 250 mg TYROSINE 20 mg

TRYPTOPHAN 90 mg

mEq/L

ACETATE 42 CHLORIDE 20

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

**STERILE** 

SINGLE DOSE CONTAINER

ROOM TEMPERATURE (25°C/77°F) AVOID EXCESSIVE HEAT PROTECT FROM FREEZING SEE PRESCRIBING INFORMATION

### Baxter

BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA

LOT

**EXP** 

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0

2B7732 NDC 0338-1139-03 **CLINIMIX** 5/25

0

0

SULFITE-FREE

(5% Amino Acids in 25% Dextrose) Injection

### **CENTRAL LINE INFUSION ONLY**

500 mL INJECTION PORT CHAMBER 50% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 10% Amino Acid Injection

Rx Only

0

0

**ACTIVATE SEAL AND** MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED DEXTROSE HYDROUS USP

ESSENTIAL AMINO ACIDS 365 mg LEUCINE ISOLEUCINE 300 mg VALINE 290 mg

LYSINE (ADDED AS THE HYDROCHLORIDE SALT) 290 mg PHENYLALANINE 290 mg 240 mg 210 mg THREONINE METHIONINE 200 mg TRYPTOPHAN 90 mg

NONESSENTIAL AMINO ACIDS

ALANINE 1035 mg ARGININE 575 mg GLYCINE 515 mg PROLINE 340 mg 250 mg SERINE 20 mg TYROSINE

mEq/L ACETATE CHLORIDE

BALANCED BY IONS FROM AMINO ACIDS PH ADJUSTED WITH GLACIAL ACETIC ACID

STERILE SINGLE DOSE CONTAINER

Baxter

BAXTER HEALTHCARE CORPORATION DEERFIELD L 60016 USA

**Container Label** 

**LOT EXP** 

2B7732 NDC 0338-1139-03

**CLINIMIX** 5/25 **SULFITE-FREE** (5% Amino Acid in 25% Dextrose) Injection

500 mL INJECTION PORT CHAMBER 50% Dextrose Injection USP

500 mL OUTLET PORT CHAMBER 10% Amino Acid Injection

Rx Only

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 1000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE

**ADMIXED INJECTION** 

DEXTROSE HYDROUS USP 25 g

#### **ESSENTIAL AMINO ACIDS**

LEUCINE 365 mg

ISOLEUCINE 300 mg

VALINE 290 mg

LYSINE (ADDED AS THE

HYDROCHLORIDE SALT) 290 mg

PHENYLALANINE 280 mg

HISTIDINE 240 mg

THREONINE 210 mg

METHIONINE 200 mg

TRYPTOPHAN 90 mg

#### NONESSENTIAL AMINO ACIDS

ALANINE 1035 mg

ARGININE 575 mg

GLYCINE 515 mg

PROLINE 340 mg

SERINE 250 mg

TYROSINE 20 mg

mEq/L

**ACETATE 42** 

CHLORIDE 20

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

**STERILE** 

SINGLE DOSE CONTAINER

**Baxter** 

BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA

LOT EXP

2B7711 NDC 0338-1103-04
CLINIMIX
5/25

## SULFITE-FREE

0

(5% Amino Acids in 25% Dextrose) Injection

### CENTRAL LINE INFUSION ONLY

1000 mL INJECTION PORT CHAMBER 50% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER

10% Amino Acid Injection

Rx Only

0

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON ACTIVATION

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION
DEXTROSE HYDROUS USP 25 g
ESSENTIAL AMINO ACIDS

 LEUCINE
 365 mg

 ISOLEUCINE
 300 mg

 VALINE
 290 mg

LYSINE (ADDED AS THE
HYDROCHLORIDE SALT)
PHENYLALANINE
HISTIDINE
THREONINE
METHIONINE
TRYPTOPHAN
200 mg
200 mg
200 mg
200 mg

NONESSENTIAL AMINO ACIDS

ALANINE 1035 mg
ARGININE 575 mg
GLYCINE 515 mg
PROLINE 340 mg
SERINE 250 mg
TYROSINE 20 mg

mEq/L

ACETATE 42 CHLORIDE 20

PH ADJUSTED WITH GLACIAL ACETIC ACID

STERILE SINGLE DOSE CONTAINER

Baxter
BAXTER HEALTHCARE CORPORATION
DEERFIELD IL 60015 USA
MADE IN USA

0

**Container Label** 

0

#### LOT EXP

0

2B7711 NDC 0338-1103-04

CLINIMIX 5/25 SULFITE-FREE (5% Amino Acid in 25% Dextrose) Injection

1000 mL INJECTION PORT CHAMBER 50% Dextrose Injection USP

1000 mL OUTLET PORT CHAMBER 10% Amino Acid Injection

Rx Only

ACTIVATE SEAL AND MIX THOROUGHLY BEFORE USE

SEE PRESCRIBING INFORMATION FOR INSTRUCTIONS ON

**ACTIVATION** 

AFTER MIXING THE PRODUCT REPRESENTS 2000 mL

REFRIGERATED STORAGE IS LIMITED TO 9 DAYS

A SLIGHT YELLOW COLOR DOES NOT ALTER THE QUALITY

AND EFFICACY OF THIS PRODUCT

ASK PHARMACIST ABOUT ADDITIVE COMPATIBILITY

CONTENTS OF EACH 100 mL OF THE ADMIXED INJECTION

DEXTROSE HYDROUS USP 25 g

### **ESSENTIAL AMINO ACIDS**

LEUCINE 365 mg

ISOLEUCINE 300 mg

VALINE 290 mg

LYSINE (ADDED AS THE

HYDROCHLORIDE SALT) 290 mg

PHENYLALANINE 280 mg

HISTIDINE 240 mg

THREONINE 210 mg

METHIONINE 200 mg

TRYPTOPHAN 90 mg

#### NONESSENTIAL AMINO ACIDS

ALANINE 1035 mg

ARGININE 575 mg

GLYCINE 515 mg

PROLINE 340 mg

SERINE 250 mg

TYROSINE 20 mg

mEq/L

**ACETATE 42** 

**CHLORIDE 20** 

BALANCED BY IONS FROM AMINO ACIDS pH ADJUSTED WITH GLACIAL ACETIC ACID

**STERILE** 

SINGLE DOSE CONTAINER

#### **Baxter**

BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA

#### **CLINIMIX**

leucine, phenylalanine, lysine, methionine, isoleucine, valine, histidine, threonine, tryptophan, alanine, glycine, arginine, proline, serine, tyrosine, dextrose injection

Product In	ıformation
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Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0338-1132

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	201 mg in 100 mL
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	154 mg in 100 mL
LYSINE (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	159 mg in 100 mL
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII: AE28 F7PNPL)	METHIONINE	110 mg in 100 mL
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	165 mg in 100 mL
VALINE (UNII: HG18 B9 YRS7) (VALINE - UNII:HG18 B9 YRS7)	VALINE	160 mg in 100 mL
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	132 mg in 100 mL
THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	116 mg in 100 mL
TRYPTO PHAN (UNII: 8 DUH1N11BX) (TRYPTO PHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	50 mg in 100 mL
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	570 mg in 100 mL
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	283 mg in 100 mL
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	316 mg in 100 mL
PROLINE (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	187 mg in 100 mL
<b>SERINE</b> (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	138 mg in 100 mL
TYROSINE (UNII: 42HK56048U) (TYROSINE - UNII:42HK56048U)	TYROSINE	11 mg in 100 mL
DEXTROSE (UNII: IY9 XDZ35W2) (DEXTROSE - UNII:IY9 XDZ35W2)	DEXTROSE	5 g in 100 mL

Inactive Ingredients			
Ingredient Name	Strength		
ACETIC ACID (UNII: Q40Q9N063P)			
WATER (UNII: 059QF0KO0R)			
NITRO GEN (UNII: N762921K75)			

Packaging				
#	Item Code	Package Description	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
1	NDC:0338-1132-03	1000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997	05/14/2018

Marketing Information				
Marketing Category Application Number or Monograph Citation Marketing Start Date Marketing E				
NDA	NDA020734	09/29/1997	05/14/2018	

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1083
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	201 mg in 100 mL	
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	154 mg in 100 mL	
LYSINE (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	159 mg in 100 mL	
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII: AE28 F7PNPL)	METHIONINE	110 mg in 100 mL	
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	165 mg in 100 mL	
VALINE (UNII: HG18B9 YRS7) (VALINE - UNII:HG18B9 YRS7)	VALINE	160 mg in 100 mL	
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	132 mg in 100 mL	
THREO NINE (UNII: 2ZD004190S) (THREONINE - UNII:2ZD004190S)	THREONINE	116 mg in 100 mL	
TRYPTOPHAN (UNII: 8 DUH1N11BX) (TRYPTOPHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	50 mg in 100 mL	
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	570 mg in 100 mL	
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	283 mg in 100 mL	
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	316 mg in 100 mL	
PROLINE (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	187 mg in 100 mL	
<b>SERINE</b> (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	138 mg in 100 mL	
TYRO SINE (UNII: 42HK56048U) (TYRO SINE - UNII:42HK56048U)	TYROSINE	11 mg in 100 mL	
DEXTROSE (UNII: IY9 XDZ35W2) (DEXTROSE - UNII:IY9 XDZ35W2)	DEXTROSE	5 g in 100 mL	

Inactive Ingredients	
Ingredient Name	Strength
ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
NITRO GEN (UNII: N762921K75)	

Packaging				
# Item Code Package Description Marketing Start Date Marketing End I				
1	NDC:0338-1083-04	2000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997	05/14/2018

Marketing Information				
Marketing Category Application Number or Monograph Citation Marketing Start Date Marketing End				
NDA	NDA020734	09/29/1997	05/14/2018	

leucine, phenylalanine, lysine, methionine, isoleucine, valine, histidine, threonine, tryptophan, alanine, glycine, arginine, proline, serine, tyrosine, dextrose injection

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1133
Route of Administration	INTRAVENOUS		

## **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	311 mg in 100 mL
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	238 mg in 100 mL
<b>LYSINE</b> (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	247 mg in 100 mL
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII: AE28 F7PNPL)	METHIO NINE	170 mg in 100 mL
ISO LEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	255 mg in 100 mL
VALINE (UNII: HG18 B9 YRS7) (VALINE - UNII:HG18 B9 YRS7)	VALINE	247 mg in 100 mL
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	204 mg in 100 mL
THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	179 mg in 100 mL
TRYPTOPHAN (UNII: 8 DUH1N11BX) (TRYPTOPHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	77 mg in 100 mL
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	880 mg in 100 mL
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	438 mg in 100 mL
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	489 mg in 100 mL
PROLINE (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	289 mg in 100 mL
SERINE (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	213 mg in 100 mL
TYRO SINE (UNII: 42HK56048U) (TYRO SINE - UNII:42HK56048U)	TYROSINE	17 mg in 100 mL
DEXTRO SE (UNII: IY9 XDZ35W2) (DEXTRO SE - UNII:IY9 XDZ35W2)	DEXTROSE	5 g in 100 mL

Inactive Ingredients	
Ingredient Name	Strength
ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
NITRO GEN (UNII: N762921K75)	

Packaging				
#	Item Code	Package Description	<b>Marketing Start Date</b>	Marketing End Date
1 1	NDC:0338-1133-03	1000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA020734	09/29/1997	

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1089
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	311 mg in 100 mL	

PHENYLALANINE (UNII: 47E5O 17Y3R) (PHENYLALANINE - UNII:47E5O 17Y3R)	PHENYLALANINE	238 mg in 100 mL
<b>LYSINE</b> (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	247 mg in 100 mL
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII: AE28 F7PNPL)	METHIO NINE	170 mg in 100 mL
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	255 mg in 100 mL
VALINE (UNII: HG18 B9 YRS7) (VALINE - UNII:HG18 B9 YRS7)	VALINE	247 mg in 100 mL
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	204 mg in 100 mL
THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	179 mg in 100 mL
TRYPTO PHAN (UNII: 8 DUH1N11BX) (TRYPTO PHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	77 mg in 100 mL
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	880 mg in 100 mL
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	438 mg in 100 mL
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	489 mg in 100 mL
PROLINE (UNII: 9DLQ4CIU6V) (PROLINE - UNII:9DLQ4CIU6V)	PROLINE	289 mg in 100 mL
<b>SERINE</b> (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	213 mg in 100 mL
TYRO SINE (UNII: 42HK56048U) (TYRO SINE - UNII:42HK56048U)	TYROSINE	17 mg in 100 mL
DEXTROSE (UNII: IY9 XDZ35W2) (DEXTROSE - UNII:IY9 XDZ35W2)	DEXTROSE	5 g in 100 mL

Inactive Ingredients	
Ingredient Name	Strength
ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
NITRO GEN (UNII: N762921K75)	

P	Packaging				
#	Item Code	Package Description	<b>Marketing Start Date</b>	Marketing End Date	
1	NDC:0338-1089-04	2000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA020734	09/29/1997	

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1134	
Route of Administration	INTRAVENOUS			

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	311 mg in 100 mL	
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	238 mg in 100 mL	
<b>LYSINE</b> (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	247 mg in 100 mL	

METHIO NINE (UNII: AE28 F7PNPL) (METHIONINE - UNII:AE28 F7PNPL)	METHIO NINE	170 mg in 100 mL
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	255 mg in 100 mL
VALINE (UNII: HG18 B9 YRS7) (VALINE - UNII:HG18 B9 YRS7)	VALINE	247 mg in 100 mL
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	204 mg in 100 mL
THREO NINE (UNII: 2ZD004190S) (THREONINE - UNII:2ZD004190S)	THREONINE	179 mg in 100 mL
TRYPTOPHAN (UNII: 8 DUH1N11BX) (TRYPTOPHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	77 mg in 100 mL
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	880 mg in 100 mL
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	438 mg in 100 mL
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	489 mg in 100 mL
PROLINE (UNII: 9DLQ4CIU6V) (PROLINE - UNII:9DLQ4CIU6V)	PROLINE	289 mg in 100 mL
SERINE (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	213 mg in 100 mL
TYROSINE (UNII: 42HK56048U) (TYROSINE - UNII:42HK56048U)	TYROSINE	17 mg in 100 mL
DEXTROSE (UNII: IY9 XDZ35W2) (DEXTROSE - UNII:IY9 XDZ35W2)	DEXTROSE	10 g in 100 mL

Inactive Ingredients	
Ingredient Name	Strength
ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
NITRO GEN (UNII: N762921K75)	

Packaging					
#	Item Code	Package Description	<b>Marketing Start Date</b>	<b>Marketing End Date</b>	
1	NDC:0338-1134-03	1000 mL in 1 BAG; Type 0: Not a Combination Product	03/20/2012		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA020734	03/20/2012	

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1091
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	311 mg in 100 mL	
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	238 mg in 100 mL	
<b>LYSINE</b> (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	247 mg in 100 mL	
METHIO NINE (UNII: AE28 F7 PNPL) (METHIO NINE - UNII: AE28 F7 PNPL)	METHIO NINE	170 mg in 100 mL	
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	255 mg in 100 mL	

VALINE (UNII: HG18B9YRS7) (VALINE - UNII:HG18B9YRS7)	VALINE	247 mg in 100 mL
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	204 mg in 100 mL
THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	179 mg in 100 mL
TRYPTOPHAN (UNII: 8 DUH1N11BX) (TRYPTOPHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	77 mg in 100 mL
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	880 mg in 100 mL
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	438 mg in 100 mL
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	489 mg in 100 mL
PROLINE (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	289 mg in 100 mL
SERINE (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	213 mg in 100 mL
TYRO SINE (UNII: 42HK56048U) (TYRO SINE - UNII:42HK56048U)	TYROSINE	17 mg in 100 mL
DEXTROSE (UNII: IY9 XDZ35W2) (DEXTROSE - UNII:IY9 XDZ35W2)	DEXTROSE	10 g in 100 mL

Inactive Ingredients	
Ingredient Name	Strength
ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
NITRO GEN (UNII: N762921K75)	

Packaging				
#	Item Code	Package Description	<b>Marketing Start Date</b>	Marketing End Date
1	NDC:0338-1091-04	2000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA020734	09/29/1997	

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1135	
Route of Administration	INTRAVENOUS			

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	311 mg in 100 mL		
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	238 mg in 100 mL		
<b>LYSINE</b> (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	247 mg in 100 mL		
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII: AE28 F7PNPL)	METHIO NINE	170 mg in 100 mL		
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	255 mg in 100 mL		
VALINE (UNII: HG18B9YRS7) (VALINE - UNII:HG18B9YRS7)	VALINE	247 mg in 100 mL		
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	204 mg in 100 mL		

THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	179 mg in 100 mL
TRYPTO PHAN (UNII: 8 DUH1N11BX) (TRYPTO PHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	77 mg in 100 mL
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	438 mg in 100 mL
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	489 mg in 100 mL
PROLINE (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	289 mg in 100 mL
SERINE (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	213 mg in 100 mL
TYRO SINE (UNII: 42HK56048U) (TYRO SINE - UNII: 42HK56048U)	TYROSINE	17 mg in 100 mL
DEXTRO SE (UNII: IY9 XDZ35W2) (DEXTRO SE - UNII:IY9 XDZ35W2)	DEXTROSE	20 g in 100 mL

Inactive Ingredients	
Ingredient Name	Strength
ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
NITRO GEN (UNII: N762921K75)	

Packaging					
#	Item Code	Package Description	<b>Marketing Start Date</b>	Marketing End Date	
1	NDC:0338-1135-03	1000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997	05/14/2018	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA020734	09/29/1997	05/14/2018

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1093	
Route of Administration	INTRAVENOUS			

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	311 mg in 100 mL	
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	238 mg in 100 mL	
<b>LYSINE</b> (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	247 mg in 100 mL	
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII: AE28 F7PNPL)	METHIONINE	170 mg in 100 mL	
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	255 mg in 100 mL	
VALINE (UNII: HG18 B9 YRS7) (VALINE - UNII: HG18 B9 YRS7)	VALINE	247 mg in 100 mL	
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	204 mg in 100 mL	
THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	179 mg in 100 mL	
TRYPTOPHAN (UNII: 8 DUH1N11BX) (TRYPTOPHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	77 mg in 100 mL	
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	880 mg in 100 mL	

GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	438 mg in 100 mL
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	489 mg in 100 mL
PROLINE (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	289 mg in 100 mL
<b>SERINE</b> (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	213 mg in 100 mL
TYRO SINE (UNII: 42HK56048U) (TYRO SINE - UNII:42HK56048U)	TYROSINE	17 mg in 100 mL
DEXTRO SE (UNII: IY9 XDZ35W2) (DEXTRO SE - UNII:IY9 XDZ35W2)	DEXTROSE	20 g in 100 mL

Inactive Ingredients	
Ingredient Name	Strength
ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
NITRO GEN (UNII: N762921K75)	

Packaging				
#	Item Code	Package Description	<b>Marketing Start Date</b>	Marketing End Date
1 1	NDC:0338-1093-04	2000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997	05/14/2018

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
NDA	NDA020734	09/29/1997	05/14/2018	

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1136	
Route of Administration	INTRAVENOUS			

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	311 mg in 100 mL	
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	238 mg in 100 mL	
LYSINE (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	247 mg in 100 mL	
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII:AE28 F7PNPL)	METHIONINE	170 mg in 100 mL	
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	255 mg in 100 mL	
VALINE (UNII: HG18 B9 YRS7) (VALINE - UNII:HG18 B9 YRS7)	VALINE	247 mg in 100 mL	
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	204 mg in 100 mL	
THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	179 mg in 100 mL	
TRYPTOPHAN (UNII: 8 DUH1N11BX) (TRYPTOPHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	77 mg in 100 mL	
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	880 mg in 100 mL	
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	438 mg in 100 mL	
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	489 mg in 100 mL	

PROLINE (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	289 mg in 100 mL
<b>SERINE</b> (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	213 mg in 100 mL
TYRO SINE (UNII: 42HK56048U) (TYRO SINE - UNII:42HK56048U)	TYROSINE	17 mg in 100 mL
DEXTRO SE (UNII: IY9 XDZ35W2) (DEXTRO SE - UNII:IY9 XDZ35W2)	DEXTROSE	25 g in 100 mL

Inactive Ingredients	
Ingredient Name	Strength
ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
NITRO GEN (UNII: N762921K75)	

Packaging				
# Item Code	Package Description	<b>Marketing Start Date</b>	<b>Marketing End Date</b>	
1 NDC:0338-1136-0	1000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997	02/29/2020	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA020734	09/29/1997	02/29/2020

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1095	
Route of Administration	INTRAVENOUS			

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	311 mg in 100 mL
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	238 mg in 100 mL
LYSINE (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	247 mg in 100 mL
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII:AE28 F7PNPL)	METHIO NINE	170 mg in 100 mL
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	255 mg in 100 mL
VALINE (UNII: HG18 B9 YRS7) (VALINE - UNII:HG18 B9 YRS7)	VALINE	247 mg in 100 mL
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	204 mg in 100 mL
THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	179 mg in 100 mL
TRYPTOPHAN (UNII: 8 DUH1N11BX) (TRYPTOPHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	77 mg in 100 mL
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	880 mg in 100 mL
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	438 mg in 100 mL
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	489 mg in 100 mL
PROLINE (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	289 mg in 100 mL
<b>SERINE</b> (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	213 mg in 100 mL

TYROSINE (UNII: 42HK56048U) (TYROSINE - UNII:42HK56048U)	TYROSINE	17 mg in 100 mL
DEXTRO SE (UNII: IY9 XDZ35W2) (DEXTRO SE - UNII:IY9 XDZ35W2)	DEXTROSE	25 g in 100 mL

Inactive Ingredients	
Ingredient Name	Strength
ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
NITRO GEN (UNII: N762921K75)	

Packaging				
#	Item Code	Package Description	<b>Marketing Start Date</b>	Marketing End Date
1	NDC:0338-1095-04	2000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997	05/14/2018

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA020734	09/29/1997	05/14/2018

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1137
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	365 mg in 100 mL
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	280 mg in 100 mL
<b>LYSINE</b> (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	290 mg in 100 mL
METHIO NINE (UNII: AE28 F7PNPL) (METHIONINE - UNII: AE28 F7PNPL)	METHIONINE	200 mg in 100 mL
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	300 mg in 100 mL
VALINE (UNII: HG18 B9 YRS7) (VALINE - UNII:HG18 B9 YRS7)	VALINE	290 mg in 100 mL
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	240 mg in 100 mL
THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	210 mg in 100 mL
TRYPTOPHAN (UNII: 8 DUH1N11BX) (TRYPTOPHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	90 mg in 100 mL
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	1035 mg in 100 mL
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	515 mg in 100 mL
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	575 mg in 100 mL
PROLINE (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	340 mg in 100 mL
SERINE (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	250 mg in 100 mL
TYROSINE (UNII: 42HK56048U) (TYROSINE - UNII:42HK56048U)	TYROSINE	20 mg in 100 mL
<b>DEXTROSE</b> (UNII: IY9 XDZ35W2) (DEXTROSE - UNII:IY9 XDZ35W2)	DEXTROSE	15 g in 100 mL

Inactive Ingredients	
Ingredient Name	Strength
ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
<b>NITRO GEN</b> (UNII: N762921K75)	

Packaging					
# Item Code	Package Description	<b>Marketing Start Date</b>	<b>Marketing End Date</b>		
1 NDC:0338-1137-03	1000 mL in 1 BAG: Type 0: Not a Combination Product	09/29/1997			

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA020734	09/29/1997	

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1099
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	365 mg in 100 mL	
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	280 mg in 100 mL	
LYSINE (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	290 mg in 100 mL	
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII: AE28 F7PNPL)	METHIONINE	200 mg in 100 mL	
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	300 mg in 100 mL	
VALINE (UNII: HG18 B9 YRS7) (VALINE - UNII:HG18 B9 YRS7)	VALINE	290 mg in 100 mL	
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	240 mg in 100 mL	
THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	210 mg in 100 mL	
TRYPTO PHAN (UNII: 8 DUH1N11BX) (TRYPTO PHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	90 mg in 100 mL	
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	1035 mg in 100 mL	
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	515 mg in 100 mL	
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	575 mg in 100 mL	
PROLINE (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	340 mg in 100 mL	
SERINE (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	250 mg in 100 mL	
TYRO SINE (UNII: 42HK56048U) (TYROSINE - UNII:42HK56048U)	TYROSINE	20 mg in 100 mL	
<b>DEXTROSE</b> (UNII: IY9 XDZ35W2) (DEXTROSE - UNII:IY9 XDZ35W2)	DEXTROSE	15 g in 100 mL	

Inactive Ingredients			
Ingredient Name	Strength		
ACETIC ACID (UNII: Q40Q9N063P)			
WATER (UNII: 059QF0KO0R)			
NITRO GEN (UNII: N762921K75)			

Packaging					
#	Item Code	Package Description	<b>Marketing Start Date</b>	Marketing End Date	
1	NDC:0338-1099-04	2000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA020734	09/29/1997	

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1138
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	365 mg in 100 mL	
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	280 mg in 100 mL	
<b>LYSINE</b> (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	290 mg in 100 mL	
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII: AE28 F7PNPL)	METHIONINE	200 mg in 100 mL	
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	300 mg in 100 mL	
VALINE (UNII: HG18B9 YRS7) (VALINE - UNII: HG18B9 YRS7)	VALINE	290 mg in 100 mL	
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	240 mg in 100 mL	
THREO NINE (UNII: 2ZD004190S) (THREONINE - UNII:2ZD004190S)	THREONINE	210 mg in 100 mL	
TRYPTOPHAN (UNII: 8 DUHIN11BX) (TRYPTOPHAN - UNII:8 DUHIN11BX)	TRYPTOPHAN	90 mg in 100 mL	
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	1035 mg in 100 mL	
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	515 mg in 100 mL	
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	575 mg in 100 mL	
<b>PROLINE</b> (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	340 mg in 100 mL	
SERINE (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	250 mg in 100 mL	
TYROSINE (UNII: 42HK56048U) (TYROSINE - UNII:42HK56048U)	TYROSINE	20 mg in 100 mL	
<b>DEXTRO SE</b> (UNII: IY9 XDZ35W2) (DEXTRO SE - UNII:IY9 XDZ35W2)	DEXTROSE	20 g in 100 mL	

Inactive Ingredients			
Ingredient Name	Strength		

ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
NITRO GEN (UNII: N762921K75)	

Packaging				
# Item Code	Package Description	<b>Marketing Start Date</b>	Marketing End Date	
1 NDC:0338-1138-03	1000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
NDA	NDA020734	09/29/1997		

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1101	
Route of Administration	INTRAVENOUS			

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	365 mg in 100 mL		
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	280 mg in 100 mL		
<b>LYSINE</b> (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	290 mg in 100 mL		
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII: AE28 F7PNPL)	METHIO NINE	200 mg in 100 mL		
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	300 mg in 100 mL		
VALINE (UNII: HG18B9YRS7) (VALINE - UNII:HG18B9YRS7)	VALINE	290 mg in 100 mL		
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	240 mg in 100 mL		
THREO NINE (UNII: 2ZD004190S) (THREONINE - UNII:2ZD004190S)	THREONINE	210 mg in 100 mL		
TRYPTO PHAN (UNII: 8 DUH1N11BX) (TRYPTO PHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	90 mg in 100 mL		
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	1035 mg in 100 mL		
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	515 mg in 100 mL		
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	575 mg in 100 mL		
<b>PROLINE</b> (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	340 mg in 100 mL		
SERINE (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	250 mg in 100 mL		
TYRO SINE (UNII: 42HK56048U) (TYRO SINE - UNII:42HK56048U)	TYROSINE	20 mg in 100 mL		
<b>DEXTRO SE</b> (UNII: IY9 XDZ35W2) (DEXTRO SE - UNII:IY9 XDZ35W2)	DEXTROSE	20 g in 100 mL		

Inactive Ingredients			
Ingredient Name Strength			
ACETIC ACID (UNII: Q40Q9N063P)			
WATER (UNII: 059QF0KO0R)			

## Packaging

	#	Item Code	Package Description	<b>Marketing Start Date</b>	Marketing End Date
l	1	NDC:0338-1101-04	2000 mL in 1 BAG: Type 0: Not a Combination Product	09/29/1997	

## **Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
NDA	NDA020734	09/29/1997		

## **CLINIMIX**

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1139	
Route of Administration	INTRAVENOUS			

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	365 mg in 100 mL		
PHENYLALANINE (UNII: 47E5O17Y3R) (PHENYLALANINE - UNII:47E5O17Y3R)	PHENYLALANINE	280 mg in 100 mL		
<b>LYSINE</b> (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	290 mg in 100 mL		
METHIO NINE (UNII: AE28 F7PNPL) (METHIO NINE - UNII: AE28 F7PNPL)	METHIO NINE	200 mg in 100 mL		
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	300 mg in 100 mL		
VALINE (UNII: HG18 B9 YRS7) (VALINE - UNII:HG18 B9 YRS7)	VALINE	290 mg in 100 mL		
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	240 mg in 100 mL		
THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	210 mg in 100 mL		
TRYPTOPHAN (UNII: 8 DUH1N11BX) (TRYPTOPHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	90 mg in 100 mL		
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	1035 mg in 100 mL		
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	515 mg in 100 mL		
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	575 mg in 100 mL		
<b>PROLINE</b> (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	340 mg in 100 mL		
<b>SERINE</b> (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	250 mg in 100 mL		
TYROSINE (UNII: 42HK56048U) (TYROSINE - UNII:42HK56048U)	TYROSINE	20 mg in 100 mL		
<b>DEXTROSE</b> (UNII: IY9 XDZ35W2) (DEXTROSE - UNII:IY9 XDZ35W2)	DEXTROSE	25 g in 100 mL		

Inactive Ingredients	
Ingredient Name	Strength
ACETIC ACID (UNII: Q40Q9N063P)	
WATER (UNII: 059QF0KO0R)	
NITRO GEN (UNII: N762921K75)	

Packaging				
#	Item Code	Package Description	<b>Marketing Start Date</b>	Marketing End Date
1	NDC:0338-1139-03	1000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997	02/29/2020

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
NDA	NDA020734	09/29/1997	02/29/2020	

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0338-1103	
Route of Administration	INTRAVENOUS			

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
LEUCINE (UNII: GMW67QNF9C) (LEUCINE - UNII:GMW67QNF9C)	LEUCINE	365 mg in 100 mL	
<b>LYSINE</b> (UNII: K3Z4F929H6) (LYSINE - UNII:K3Z4F929H6)	LYSINE	290 mg in 100 mL	
METHIO NINE (UNII: AE28 F7PNPL) (METHIONINE - UNII: AE28 F7PNPL)	METHIONINE	200 mg in 100 mL	
ISOLEUCINE (UNII: 04Y7590D77) (ISOLEUCINE - UNII:04Y7590D77)	ISOLEUCINE	300 mg in 100 mL	
VALINE (UNII: HG18B9YRS7) (VALINE - UNII:HG18B9YRS7)	VALINE	290 mg in 100 mL	
HISTIDINE (UNII: 4QD397987E) (HISTIDINE - UNII:4QD397987E)	HISTIDINE	240 mg in 100 mL	
THREO NINE (UNII: 2ZD004190S) (THREO NINE - UNII:2ZD004190S)	THREONINE	210 mg in 100 mL	
TRYPTO PHAN (UNII: 8 DUH1N11BX) (TRYPTO PHAN - UNII:8 DUH1N11BX)	TRYPTOPHAN	90 mg in 100 mL	
ALANINE (UNII: OF5P57N2ZX) (ALANINE - UNII:OF5P57N2ZX)	ALANINE	1035 mg in 100 mL	
GLYCINE (UNII: TE7660 XO1C) (GLYCINE - UNII:TE7660 XO1C)	GLYCINE	515 mg in 100 mL	
ARGININE (UNII: 94ZLA3W45F) (ARGININE - UNII:94ZLA3W45F)	ARGININE	575 mg in 100 mL	
<b>PROLINE</b> (UNII: 9 DLQ4CIU6 V) (PROLINE - UNII:9 DLQ4CIU6 V)	PROLINE	340 mg in 100 mL	
SERINE (UNII: 452VLY9402) (SERINE - UNII:452VLY9402)	SERINE	250 mg in 100 mL	
TYROSINE (UNII: 42HK56048U) (TYROSINE - UNII:42HK56048U)	TYROSINE	20 mg in 100 mL	
DEXTRO SE (UNII: IY9 XDZ35W2) (DEXTRO SE - UNII:IY9 XDZ35W2)	DEXTROSE	25 g in 100 mL	

Inactive Ingredients			
Ingredient Name	Strength		
ACETIC ACID (UNII: Q40Q9N063P)			
WATER (UNII: 059QF0KO0R)			
NITRO GEN (UNII: N762921K75)			

P	ackaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date

1 NDC:0338-1103-04	2000 mL in 1 BAG; Type 0: Not a Combination Product	09/29/1997	05/14/2018
Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA020734	09/29/1997	05/14/2018

# Labeler - Baxter Healthcare Corporation (005083209)

Establishment			
Name	Address	ID/FEI	Business Operations
Baxter Healthcare Corporation		189326168	ANALYSIS(0338-1132, 0338-1083, 0338-1133, 0338-1089, 0338-1134, 0338-1091, 0338-1135, 0338-1093, 0338-1136, 0338-1095, 0338-1137, 0338-1099, 0338-1138, 0338-1101, 0338-1139, 0338-1103), MANUFACTURE(0338-1132, 0338-1083, 0338-1133, 0338-1089, 0338-1134, 0338-1091, 0338-1135, 0338-1093, 0338-1136, 0338-1095, 0338-1137, 0338-1099, 0338-1138, 0338-1101, 0338-1139, 0338-1103), LABEL(0338-1132, 0338-1083, 0338-1133, 0338-1089, 0338-1134, 0338-1091, 0338-1135, 0338-1093, 0338-1136, 0338-1095, 0338-1137, 0338-1099, 0338-1138, 0338-1101, 0338-1139, 0338-1103), PACK(0338-1132, 0338-1083, 0338-1133, 0338-1089, 0338-1134, 0338-1091, 0338-1135, 0338-1093, 0338-1136, 0338-1095, 0338-1137, 0338-1099, 0338-1138, 0338-1101, 0338-1139, 0338-1103), STERILIZE(0338-1132, 0338-1083, 0338-1133, 0338-1089, 0338-1089, 0338-1134, 0338-1091, 0338-1135, 0338-1093, 0338-1136, 0338-1095, 0338-1137, 0338-1099, 0338-1138, 0338-1101, 0338-1139, 0338-1103)

Establishment			
Name	Address	ID/FEI	Business Operations
Baxter Healthcare Corporation		059140764	ANALYSIS(0338-1132, 0338-1083, 0338-1133, 0338-1089, 0338-1134, 0338-1091, 0338-1135, 0338-1093, 0338-1136, 0338-1095, 0338-1137, 0338-1099, 0338-1138, 0338-1101, 0338-1139, 0338-1103), MANUFACTURE(0338-1132, 0338-1083, 0338-1133, 0338-1089, 0338-1134, 0338-1091, 0338-1135, 0338-1093, 0338-1136, 0338-1095, 0338-1137, 0338-1099, 0338-1138, 0338-1101, 0338-1139, 0338-1103), LABEL(0338-1132, 0338-1083, 0338-1133, 0338-1089, 0338-1134, 0338-1091, 0338-1135, 0338-1093, 0338-1136, 0338-1095, 0338-1137, 0338-1099, 0338-1138, 0338-1101, 0338-1139, 0338-1103), PACK(0338-1132, 0338-1083, 0338-1133, 0338-1089, 0338-1134, 0338-1091, 0338-1135, 0338-1093, 0338-1136, 0338-1095, 0338-1137, 0338-1099, 0338-1138, 0338-1101, 0338-1139, 0338-1103), STERILIZE(0338-1132, 0338-1095, 0338-1133, 0338-1137, 0338-1089, 0338-1134, 0338-1091, 0338-1135, 0338-1093, 0338-1136, 0338-1095, 0338-1137, 0338-1099, 0338-1138, 0338-1101, 0338-1139, 0338-1103)

Revised: 10/2018 Baxter Healthcare Corporation