HEPARIN SODIUM - heparin sodium injection, solution Cantrell Drug Company

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

Heparin Sodium 2,500 USP Units Added to 4.5% Sodium Chloride 500 mL Bag

HEPARIN Sodium 2,500 USP Units

Added to 0.45% Sodium Chloride 500 mL* Bag

(5 USP units/mL) *Volume & Concentration Exclude Manufacturer Overfill Store at Room Temperature. Single-Dose Bag. Hospital/Office Use Only. Injection Solution For IV Use.







Each mL Contains: Heparin Sodium 5 USP units, Sodium Chloride 4.54 mg, trace amount of Benzyl Alcohol. pH adj: HCl/NaOH. Outsourced Compounded Drug. Not for Resale.

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LOT: xxxxx BUD:

CMPD Date: 03/13



HEPARIN SODIUM

heparin sodium injection, solution

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:52533-096
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
Heparin Sodium (UNII: ZZ45AB24CA) (Heparin - UNII:T2410KM04A)	Heparin	5 [USP'U] in 1 mL

Inactive Ingredients			
Ingredient Name	Strength		
SODIUM CHLORIDE (UNII: 451W47IQ8X)	4.5 mg in 1 mL		
BENZYL ALCOHOL (UNII: LKG8494WBH)	0.00005 mL in 1 mL		
Water (UNII: 059QF0KO0R)			

Other Ingredients			
Ingredient Kind	Ingredient Name	Quantity	
May contain	HYDRO CHLO RIC ACID (UNII: QTT17582CB)		
May contain	SO DIUM HYDRO XIDE (UNII: 55X04QC32I)		

Packaging			
# Item Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:52533-096-40	500 mL in 1 BAG		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		02/18/2011	

Labeler - Cantrell Drug Company (035545763)

Revised: 12/2014 Cantrell Drug Company