## OXYTOCIN - oxytocin 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.

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Oxytocin 30 USP Units Added to 0.9% Sodium Chloride 500 mL Bag



Store at Room Temperature. Preservative Free. Single-Dose Bag. Injection Solution for IV Use Only.

Volume: 500 mL\*

**Rx Only** 

\*Volume Excludes Manufacturer Overfill

Each 500 mL Bag Contains: Oxytocin 30 USP Units added to 0.9%

Sodium Chloride. pH adj: Glacial Acetic Acid/Sodium Hydroxide.



NDC: 52533-056-30

Lot: xxxxx

BUD:

**CMPD Date: 03/13** 

00001

Hospital/Office Use Only
Outsourced Compounded Drug. Not for Resale.





Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:52533-056
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
OXYTOCIN (UNII: 1JQS135EYN) (OXYTOCIN - UNII:1JQS135EYN)	OXYTOCIN	30 [USP'U] in 500 mL	

Inactive Ingredients			
Ingredient Name	Strength		
Sodium Chloride (UNII: 451W47IQ8X)	4.5 g in 500 mL		
Water (UNII: 059QF0KO0R)			

Other Ingredients			
Ingredient Kind	Ingredient Name	Quantity	
May contain	ACETIC ACID (UNII: Q40Q9N063P)		
May contain	SODIUM HYDRO XIDE (UNII: 55X04QC32I)		

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:52533-056-30	500 mL in 1 BAG			

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		10/20/2011	

## Labeler - Cantrell Drug Company (035545763)

Revised: 12/2014 Cantrell Drug Company