QUALITY CHOICE IODINE TINCTURE MILD- iodine and sodium iodide and alcohol liquid Chain Drug Market Association

Quality Choice Iodine Ticture

Drug Facts

Active ingredient

Iodine Tincture, USP 2%, Mild

Purpose

First Aid Antiseptic

Indications

First aid to help prevent infection in minor cuts, scrapes and burns

Warnings

For external use only.

Ask a doctor before use if you have

- deep or puncture wounds
- animal bites
- serious burns
- Flammable: Keep away from sparks heat and flame

Stop use and consult doctor if

• the condition persists or gets worse, or if using for longer than one week

When using this product

- do not use in the eyes. If contact occurs, flush with large amounts of water while lifting upper and lower lids
- do not apply over large areas of the body

Keep out of reach of children.

In case of accidental ingestion, give milk then a starch solution made by mixing two tablespoonfuls of cornstarch or flour to a pint of water. Contact a Poison Control Center immediately.

Directions

- clean the affected area
- apply a small amount on the area 1 to 3 times daily
- may be covered with sterile bandage
- if bandaged let dry first

Other information

• will stain skin and clothing

Inactive ingredient

none

Principal Display Panel



QUALITY CHOICE IODINE TINCTURE MILD

iodine and sodium iodide and alcohol liquid

Product Information							
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:63868-330			
Route of Administration	TOPICAL						
Active Increalizat/Active	Maiaha						
Active Ingredient/Active Moiety							
Ingredient Name			Basis of Strengt	th Strength			
IODINE (UNII: 9679TC07X4) (IODINE - UNII:9679TC07X4)			IODINE	20 mg in 1 mL			
SODIUM IODIDE (UNII: F5WR8N145C) (IODIDE ION - UNII:09G4I6V86Q)			IODIDE ION	20.4 mg in 1 mL			
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)			ALCOHOL	470 mg in 1 mL			
Inactive Ingredients							

Ingredient Name			Strength	
WATER (UNII: 059QF	F0KO0R)			
Packaging				
# Item Code	Package Description	Marketing Start Date	Marketing End Date	
	30 mL in 1 BOTTLE; Type 0: Not a Combination Product	12/12/2017		
Marketing I	nformation			
Marketing	Application Number or Monograph	Marketing Start	Marketing End	
Category	Citation	Date	Date	
OTC Monograph Dru	g M003	01/01/1979		

Labeler - Chain Drug Market Association (011920774)

Registrant - Pharma Nobis, LLC (118564114)

Establishment

Name	Address	ID/FEI	Business Operations
Pharma Nobis, LLC		118564114	manufacture(63868-330),analysis(63868-330),pack(63868-330), label(63868-330)

Revised: 12/2023

Chain Drug Market Association