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

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**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