

POVIDONE IODINE PREP PAD- antiseptic swab
Dynarex Corporation

1108 Povidone Iodine Prep Pad NDC 67777-002-01
1108UB-10 Povidone Iodine Prep Pad NDC 67777-002-03

Active Ingredient

Povidone-Iodine USP 10% w/v (9.85% w/w)

Purpose

First Aid Antiseptic

Use(s)

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

Warnings

For External Use Only

Do not use

- In the eyes or over large areas of the body
- Longer than one week unless directed by a doctor

Ask a doctor before use if you have

- Deep or puncture wounds
- Animal bites
- Serious burns

Stop use and ask a doctor if

- Condition persists or gets worse

Keep out of reach of children

If swallowed, get medical help or contact a Poison Control Center (1-800-222-1222) right away.

Directions

- Tear at notch
- Remove prep pad
- Apply a small amount of this product on the area 1 to 3 times daily.

- May be covered with a sterile bandage
- If bandaged, let dry first
- Use only once

Other Information

- Store at room temperature 15º-30ºC (59º-86ºF)
- Flammable. Keep away from fire or flame

Inactive Ingredients

Anhydrous Citric Acid, Glycerin, Polysorbate 80, Sodium Citrate, Sodium Phosphate Dibasic, Water

Label

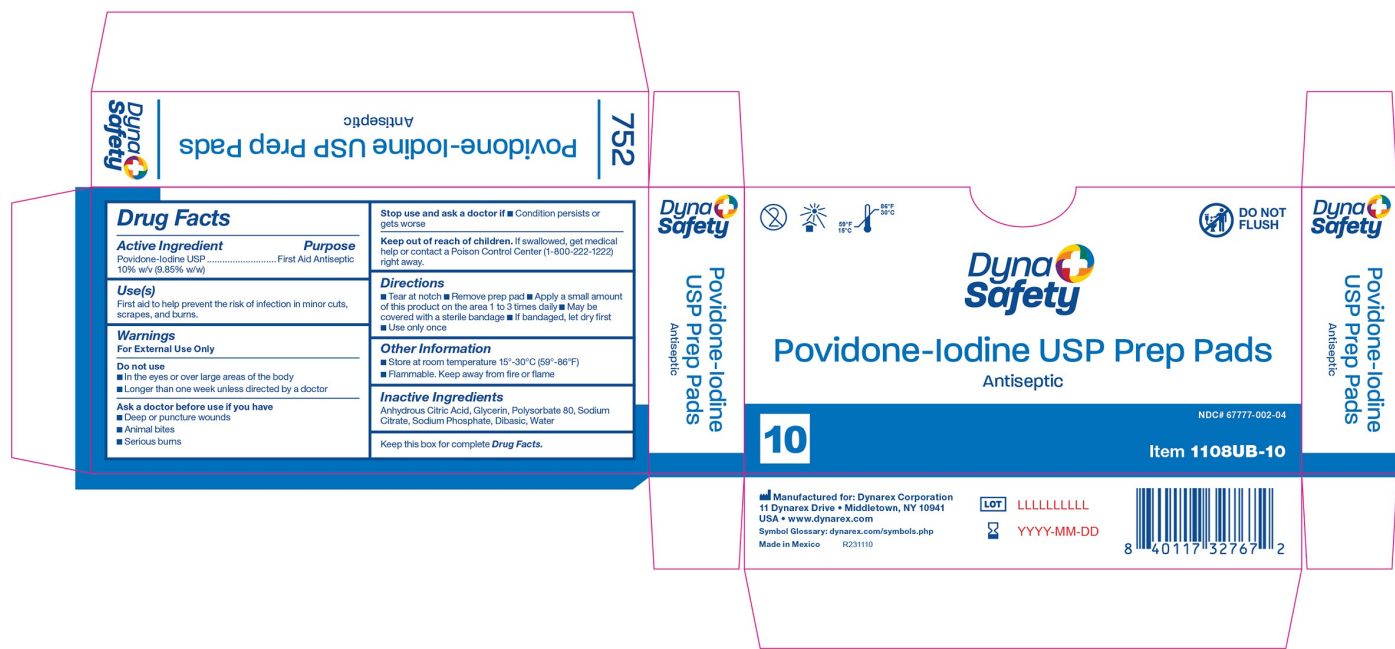


Label



Povidon Iodine Prep Pad

Label 1108UB-10



1108UB-10

POVIDONE IODINE PREP PAD			
antiseptic swab			
Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:67777-002
Route of Administration	TOPICAL		
Active Ingredient/Active Moiety			
Ingredient Name		Basis of Strength	Strength
POVIDONE-IODINE (UNII: 85H0HZU99M) (IODINE - UNII:9679TC07X4)		IODINE	10 g in 100 g
Inactive Ingredients			
Ingredient Name			Strength
GLYCERIN (UNII: PDC6A3C0OX)			
WATER (UNII: 059QF0KO0R)			
POLYSORBATE 80 (UNII: 6OZP39ZG8H)			
SODIUM CITRATE (UNII: 1Q73Q2JULR)			
SODIUM PHOSPHATE, DIBASIC (UNII: GR686LBA74)			
ANHYDROUS CITRIC ACID (UNII: XF417D3PSL)			
Product Characteristics			
Color		Score	

Shape		RECTANGLE	Size	
Flavor			Imprint Code	
Contains				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:67777-002-01	1000 in 1 CASE	12/17/2018	
1	NDC:67777-002-02	100 in 1 BOX		
1		0.3 g in 1 PACKET; Type 0: Not a Combination Product		
2	NDC:67777-002-03	1000 in 1 CASE	12/17/2018	
2	NDC:67777-002-04	10 in 1 BOX		
2		0.3 g in 1 PACKET; Type 0: Not a Combination Product		
Marketing Information				
Marketing Category		Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug		M003	12/17/2018	

Labeler - Dynarex Corporation (008124539)