VIAMED ALCOHOL PREP PAD- alcohol swab Wuxi Medical Instrument Factory

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

VIAMED Alcohol Prep Pad

Drug facts

Active Ingredient Isopropyl Alcohol 70%

Purpose

Antiseptic

Use

For preparation of skin prior to injection

Warnings

For external use only. Flammable keep away from the fire or flame

Do not use

with electrocautery procedures in the eyes

Stop use

if irritation or redness develop. If your condition persists for more than 72 hours, consult a doctor.

Keep out of the reach of children

if swallowed, get medical help or contact a Poison Control Center right away.

Directions

Wipe injectio site vigorously and discard after single use.

Inactive Ingredient

Purified water

Product labeling

ViaMed Alcohol Prep Pad For external use only One pad Saturated with 70% Isopropyl Alcohol Made in China Rece International Corp Miami Lakes, FL 33014



	nt Purpose 70% v/vAntiseptic
Use For preparatio	on of skin prior to injection
Warnings For ext	ternal use only.
Flammable, keep	away from the fire or flame.
the eves Ston Lies	electrocautery procedures *in if irritation or redness ndition persists for more than a doctor. Keep out of reach of wed, get medical help or Control Center right away.
Directions Wipe discard after single	injection site vigorously and e use.
Inactive ingredi	ent purifed water
LOT 150625	20150625 [™] 20200625 TEAR HERE

VIAMED ALCOHOL Platcohol swab	REP PAD				
Product Information					
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:42947-500	
Route of Administration	TOPICAL				
Active Ingredient/Active	Moiety				
Ingredient Name			Basis of Streng	th	Strength
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)			ALCOHOL		70 g in 100 g
Inactive Ingredients					
Ing	redient Name			Strei	ngth

W	ATER (UNII: 059Q	F0KO0R)					
P	ackaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date			
1	NDC:42947-500- 01	100 in 1 PACKAGE	09/10/2015				
1		0.36 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			
OT fin	FC monograph not al	part333E	09/10/2015				

Labeler - Wuxi Medical Instrument Factory (421292863)

Registrant - Wuxi Medical Instrument Factory (421292863)

Revised: 2/2023

Wuxi Medical Instrument Factory