CERTUS WASH TOWELETTE - benzalkonium chloride swab Certus Medical, Inc.

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.

Certus Wash Towelette 210033 Drug Facts and Label

Drug Facts Box OTC Active Ingredient Section

Benzalkonium Chloride 0.13% w/w

Drug Facts Box OTC Purpose Section

First Aid Antiseptic

Drug Facts Box OTC Indications & Usage Section

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

Drug Facts Box OTC Warnings Section

For external use only

Drug Facts Box OTC Do Not Use Section

in the eyes or apply over large areas of the body longer than 1 week unless directed by a doctor

Drug Facts Box OTC Ask Doctor Section

in case of deep or puncture wounds, animal bites, or serious burns

Drug Facts Box OTC Stop Use Section

and ask a doctor if the condition persists or gets worse

Drug Facts Box OTC Keep Out Of Reach Of Children Section

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

Drug Facts Box OTC Dosage & Administration Section

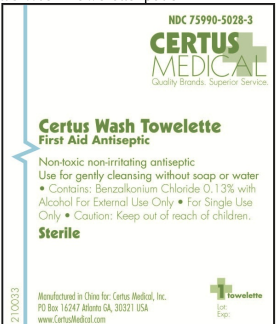
clean the affected area apply a small amount of this product on the area 1 to 3 times daily may be covered with a sterile bandage when dry

Drug Facts Box OTC Inactive Ingredient Section

fragrance, isopropyl alcohol, propylene glycol, purified water, sodium bicarbonate

Certus Wash Towelette 210033 pouch

210033.jpg Certus Wash Towelette pouch





ABW 3-23

CERTUS WASH TOWELETTE

benzalkonium chloride swab

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:75990-5028	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6JUD5X6Y)	BENZALKONIUM CHLORIDE	0.0013 mL in 1 mL		

Inactive Ingredients			
Ingredient Name	Strength		
WATER (UNII: 059QF0KO0R)			
ISOPROPYL ALCOHOL (UNII: ND2M416302)			
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)			
SODIUM BICARBONATE (UNII: 8MDF5V39QO)			

Pack	aging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date

1 NDC:75990-5028-3	1.4 mL in 1 POUCH				
Marketing Information					
Marketing Category	Application Number or Monogr	aph Citation Ma	arketing Start Date	Marketing End Date	
OTC monograph not final	part333A	05/	/0 1/20 11		

Labeler - Certus Medical, Inc. (966433653)

Revised: 4/2011 Certus Medical, Inc.