ALL STOP MEDICATED BODY WASH- benzalkonium chloride soap Q-Based Solutions 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.

All Stop Medicated Body Wash 6265 Drug Facts and Label

Drug Facts Box OTC-Active Ingredient Section

Benzalkonium Chloride 2.5%

Drug Facts Box OTC-Purpose Section

Antiseptic

Drug Facts Box OTC-Indications & Usage Section

for hand-washing to decrease bacteria on the skin

Drug Facts Box OTC-Warnings Section

For external use only

Drug Facts Box OTC-When Using Section

do not get into eyes

if contact occurs, rinse eyes thoroughly with water

Drug Facts Box OTC-Stop Use Section

irritation and redness develop

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

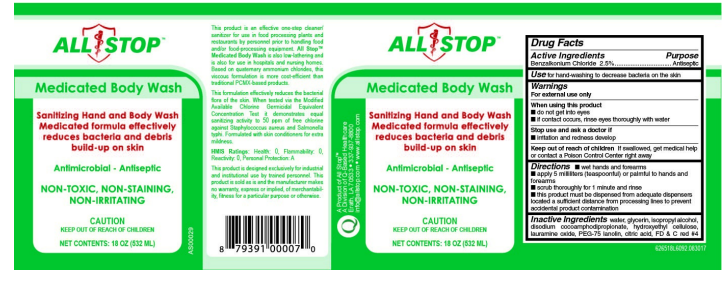
- wet hands and forearms
- apply 5 milliliters (teaspoonful) or palmful to hands and forearms
- scrub thoroughly for 1 minute and rinse

Drug Facts Box OTC-Inactive Ingredient Section

ater, glycerin, isopropyl alcohol, disodium cocoamphodiproprionate, hydroxyethylcellulose, lauramine oxide,

PEG-75 lanolin, citric acid, FD and C red no.4

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ALL STOP MEDICATED BODY WASH

benzalkonium chloride soap

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:42355-265
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
BENZALKO NIUM CHLO RIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6JUD5X6Y)	BENZALKONIUM CHLORIDE	25.0 mg in 1 mL

Inactive Ingredients				
Ingredient Name	Strength			
WATER (UNII: 059QF0KO0R)				
ISOPROPYL ALCOHOL (UNII: ND2M416302)				
HYDRO XYETHYL CELLULO SE (1500 MPA.S AT 1%) (UNII: L605B5892V)				
DISO DIUM CO CO AMPHO DIPRO PIO NATE (UNII: 6 K8 PRP397M)				
LAURAMINE O XIDE (UNII: 4F6 FC4MI8 W)				
PEG-75 LANOLIN (UNII: 09179OX7TB)				
GLYCERIN (UNII: PDC6A3C0OX)				
CITRIC ACID MONO HYDRATE (UNII: 2968 PHW8 QP)				
FD&C RED NO. 4 (UNII: X3W0 AM1JLX)				

P	Packaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:42355-265- 05	532 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	12/21/2017	

2	NDC:42355-265- 14	3785 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	12/21/2017	
3	NDC:42355-265- 19	18900 mL in 1 CONTAINER; Type 0: Not a Combination Produc	et 12/21/2017	
Marketing Information				
	Marketing Categ	ory Application Number or Monograph Citation M	Marketing Start Date	Marketing End Date
0	TC monograph not	final part333E 12	/21/2017	

Labeler - Q-Based Solutions Inc (153509315)

Registrant - ABC Compounding Co., Inc. (003284353)

Establishment				
Name	Address	ID/FEI	Business Operations	
ABC Compounding Co., Inc.		003284353	manufacture(42355-265)	

Revised: 7/2019 Q-Based Solutions Inc