FREE HAND- alcohol gel Atco International

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.

Free Hand 6605 Drug Facts and Label

Drug Facts Box OTC-Active Ingredient Section

Ethyl Alcohol 62%

Drug Facts Box OTC-Purpose Section

Antiseptic

Drug Facts Box OTC-Indications & Usage Section

for hand-washing to decrease bacteria on the skin, only when water is not available

Drug Facts Box OTC-Warnings Section

FLAMMABLE, keep away from fire and flames

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 thoroughly with product and allow to dry without wiping

Drug Facts Box OTC-Inactive Ingredient Section

water, diisopropylamine, carbomer, propylene glycol, DMDM hydantoin, tocopheryl acetate, aloe barbadensis

Free Hand 6605 18oz

Free Hand 18oz

FREE HAND

alcohol gel

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:62712-221
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	0.7 mL in 1 mL	

Inactive Ingredients			
Ingredient Name	Strength		
WATER (UNII: 059QF0KO0R)			
CARBOMER 934 (UNII: Z135WT9208)			
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)			
DMDM HYDANTO IN (UNII: BYR0546 TOW)			
ALOE VERA LEAF (UNII: ZY8 1Z8 3H0 X)			
.ALPHATO COPHEROL ACETATE, DL- (UNII: WR1WPI7EW8)			
DIISOPROPYLAMINE (UNII: BR9JLI40NO)			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:62712-221- 17	532 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	10/01/2009	
2	NDC:62712-221- 24	118 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	10/01/2009	
3	NDC:62712-221- 28	149 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	10/01/2009	

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC monograph not final	part333E	10/01/2009			

Labeler - Atco International (033504929)

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

Establishment				
Name	Address	ID/FEI	Business Operations	
ABC Compounding Co., Inc.		003284353	manufacture(62712-221)	

Revised: 12/2018 Atco International