### CAREONE TAHITIAN COCONUT- ethyl alcohol liquid AMERICAN SALES COMPANY

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.

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## **DRUG FACTS**

## **ACTIVE INGREDIENT**

ETHYL ALCOHOL 65%

## PURPOSE

ANTISEPTIC

## USES

TO DECREASE BACTERIA ON THE SKIN

## WARNINGS

FOR EXTERNAL USE ONLY. FLAMMABLE. KEEP AWAY FROM SOURCE OF HEAT OR FIRE

## WHEN USING THIS PRODUCT

AVOID CONTACT WITH EYES. IF CONTACT OCCURS, RINSE WITH WATER

STOP USE AND ASK A DOCTOR IF

IRRITATION OR REDNESS DEVELOPS AND LASTS

KEEP OUT OF REACH OF CHILDREN

IN CASE OF ACCIDENTAL INGESTION, GET MEDICAL HELP OR CONTACT A POISON CONTROL CENTER IMMEDIATELY

### DIRECTIONS

APPLY A SMALL AMOUNT TO YOUR PALM AND RUB HANDS TOGETHER BRISKLY UNTIL DRY. CHILDREN UNDER 6 YEAR OLD SHOULD BE SUPERVISED WHEN USING THIS PRODUCT

## **OTHER INFORMATION**

STORE AT A TEMPERATURE BELOW 110°F (43°C)

## **INACTIVE INGREDIENTS**

WATER, FRAGRANCE (PARFUM), CARBOMER, GLYCERIN, ISOPROPYL ALCOHOL, PROPYLENE GLYCOL, MANNITOL, CELLULOSE, HYDROXYPROPYL METHYLCELLULOSE, BENZOPHENONE-4, AMINOMETHYL PROPANOL, TOCOPHERYL ACETATE, IRON OXIDES (CI 77491, CI 77492, CI 77499), RED 4 (CI 14700), RED 33 (CI 17200), YELLOW 5 (CI 19140)



CAREONE TAHITIAN	COCONUT				
ethyl alcohol liquid					
Product Information					
Product T ype	HUMAN OTC DRUG	Item Code	(Source)	NDO	2:41520-401
Route of Administration	TOPICAL				
Active Ingredient/Active Moi	ety				
Ingre	dient Name		Basis of Strengt	h	Strength
ALCOHOL (UNII: 3K9958V90M) (ALC	COHOL - UNII:3K9958V90M)		ALCOHOL		650 mg in 1 mL
Inactive Ingredients					

		Ingredient Name				Strength	
WATI	E <b>R</b> (UNII: 059QF0KO	0 R)					
CARB	OMER 934 (UNII: Z1	35WT9208)					
GLYC	C <b>ERIN</b> (UNII: PDC6A3	COOX)					
ISOP	ROPYL ALCOHOL	(UNII: ND2M416302)					
PROF	YLENE GLYCOL (U	JNII: 6DC9Q167V3)					
MANN	NITOL (UNII: 30WL5	3L36A)					
POW	DERED CELLULOSI	E (UNII: SMD1X3XO9M)					
HYPR	OMELLOSES (UNII:	3NXW29V3WO)					
SULIS	SOBENZONE (UNII:	1W6L629B4K)					
AMIN	O METHYL PRO PAN	<b>OL</b> (UNII: LU49E6626Q)					
.ALPI	HATOCOPHEROL	ACETATE (UNII: 9E8X80D2L0)					
FERR	FERRIC OXIDE RED (UNII: 1K09F3G675)						
FERR	FERRIC OXIDE YELLOW (UNII: EX438O2MRT)						
FERROSOFERRIC OXIDE (UNII: XM0 M87F357)							
FD&C	C RED NO.4 (UNII: X	3W0 AM1JLX)					
D&C	RED NO. 33 (UNII: 9)	DBA0SBB0L)					
FD&C	<b>YELLOW NO. 5</b> (U	NII: I753WB2F1M)					
Pack	aging						
#	Item Code	Package Description	Marketing Start Date Mar		rketing End Date		
1 ND	C:41520-401-02	59 mL in 1 BOTTLE, PLASTIC					
Marketing Information							
Ma	rketing Category	Application Number or Monogra	ph Citation	Marketing Start D	Date	Marketing End Date	
OTC r	nonograph not final	part333E		02/06/2014			
		-					

# Labeler - American Sales Company (809183973)

**Registrant** - APOLLO HEALTH AND BEAUTY CARE (201901209)

# Establishment

Name	Address	ID/FEI	<b>Business Operations</b>
APOLLO HEALTH AND BEAUTY CARE		201901209	manufacture(41520-401)

Revised: 2/2014

#### AMERICAN SALES COMPANY