# ADOYA SANITIZING WIPES- alcohol cloth Jokki Labs LLC

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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### **Adoya Sanitizing Wipes**

### **Drug Facts**

### Active ingredient

Ethyl Alcohol 70%

### **Purpose**

Antiseptic

### Use

For hand washing to decrease bacteria on the skin.

### **Warnings**

For external use only.

Flammable, keep away from fire or flame.

### Do not use

in the eyes

### Stop ue and ask a doctor if

- irritation and redness develop
- condition persists for more than 72 hours

### Keep out of reach of children.

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

### **Directions**

Wet hands thoroughly with product and allow to dry without wiping.

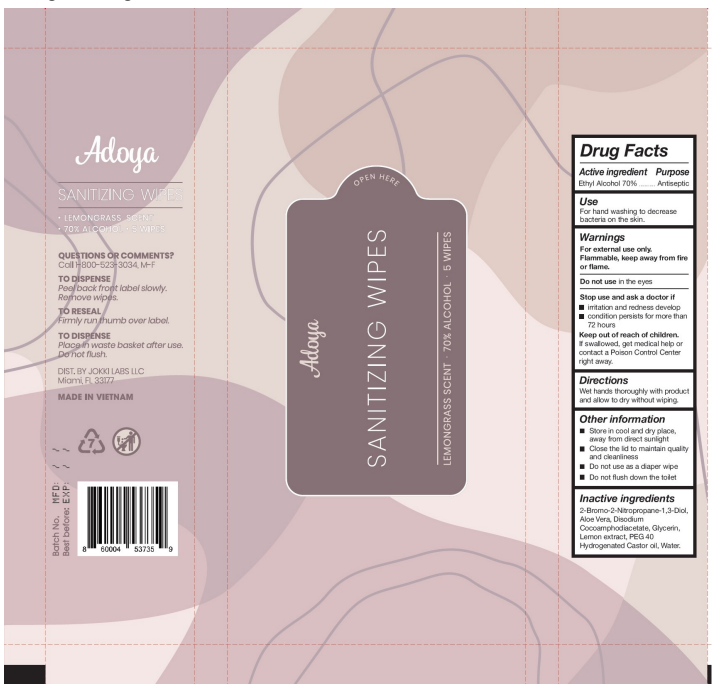
### Other information

- Store in cool and dry place, away from direct sunlight
- Close the lid to maintain quality and cleanliness
- Do not use as a diaper wipe
- Do not flush down the toilet

### **Inactive ingredients**

2-Bromo-2-Nitropropane-1,3-Diol, Aloe Vera, Disodium Cocoamphodiacetate, Glycerin, Lemon

### **Package Labeling:**



# ADOYA SANITIZING WIPES alcohol cloth Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:79708-001 Route of Administration TOPICAL Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALCOHOL (UNII: 3K9958 V90M) (ALCOHOL - UNII:3K9958 V90M)	ALCOHOL	0.7 mL in 1 mL

Inactive Ingredients		
Ingredient Name	Strength	
BRONOPOL (UNII: 6PU1E16C9W)		
ALOE VERA LEAF (UNII: ZY8 1Z8 3H0 X)		
DISO DIUM CO CO AMPHO DIACETATE (UNII: 18 L9 G3U51M)		
GLYCERIN (UNII: PDC6A3C0OX)		
LEMON (UNII: 24RS0A988O)		
POLYOXYL 40 HYDROGENATED CASTOR OIL (UNII: 7YC686GQ8F)		
WATER (UNII: 059QF0KO0R)		

l	Packaging					
	# Item Code	Package Description	Marketing Start Date	Marketing End Date		
	1 NDC:79708- 001-01	5 in 1 BAG	08/07/2020			
	1	14 mL in 1 PATCH; Type 2: Prefilled Drug Delivery Device/System (syringe, patch, etc.)				

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC monograph not final	part333E	08/07/2020			

## Labeler - Jokki Labs LLC (117570400)

Revised: 8/2020 Jokki Labs LLC