## WET WIPES- wet wipes cloth Sourcery Ltd

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

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## Alcohol-Based Wet Wipe

## Active Ingredient(s)

Alcohol, Benzalkonium Chloride, C12-14-Alkyldimethyl(Ethylbenzyl) Ammonium Chlorides, Ethylhexylglycerin, Chlorphenesin

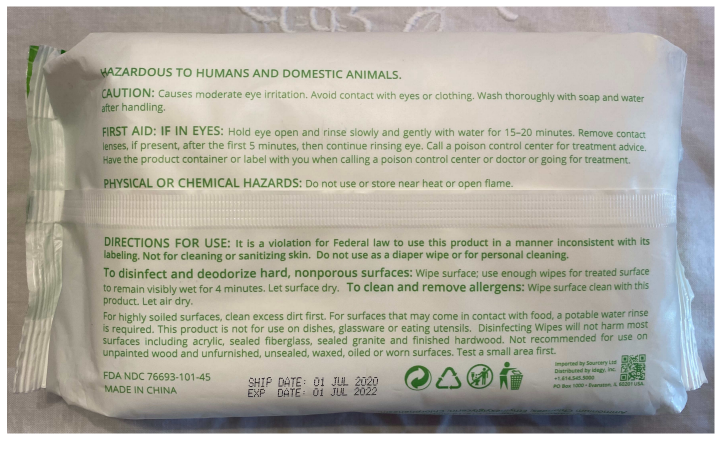
## Purpose

Antiseptic, disinfection and surface cleaning

## Use

TO disinfect and deodorize hard, nonporous surfaces.

## Warnings



#### Do not use

Do not use or store near heat or open flame.

When using this product keep out of eyes, ears, and mouth. In case of contact with eyes, rinse eyes

thoroughly with water.

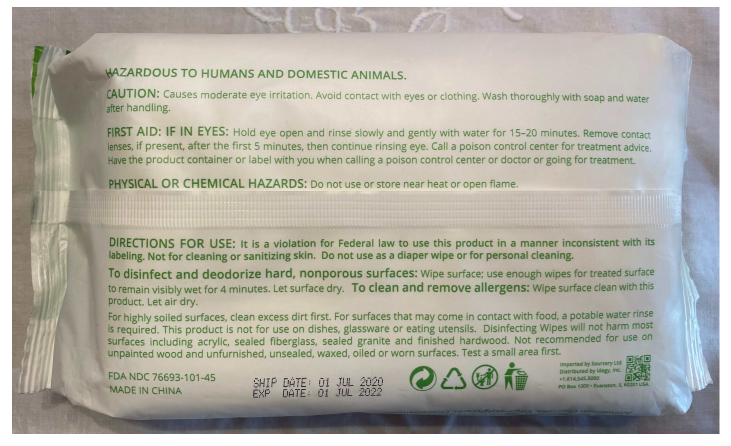
Stop use and ask a doctor if irritation or rash occurs. These may be signs of a serious condition.

Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

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## Directions



## Other information

- Store between 15-30C (59-86F)
- Avoid freezing and excessive heat above 40C (104F)
- Do not use or store near open heat or flame.

## **Inactive ingredients**

glycerin, purified water, fragrance



## Package Label - Principal Display Panel

80 wipes

NDC 7663-101-45

WET WIPES						
wet wipes cloth						
Product Information						
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:76693-101		
Route of Administration	TOPICAL					
Active Ingredient/Active Mo	iety					
Ing	Basis of Strength		Strength			
BENZALKONIUM (UNII: 7N6JUD5X6	BENZALKONIUM		0.06 g in 100 g			
PHENOXYETHANOL (UNII: HIE492Z	PHENOXYETHANOL		0.3 g in 100 g			
<b>DIDECYLDIMONIUM CHLORIDE</b> (U UNII:Z7F472XQPA)	DIDEC YL DIMO NIUM CHLO RIDE		0.4 g in 100 g			
CHLORPHENESIN (UNII: 1670 DAL4S	CHLORPHENESIN		0.1 g in 100 g			
ALCOHOL (UNII: 3K9958V90M) (AL	ALCOHOL		10 g in 100 g			
Inactive Ingredients						
Ingredient Name					Strength	

GLYCERIN (UNII: PDC6A3C0OX)						0.1 g in 100 g				
WATER (UNII: 059QF0KO0R)						48.97 g in 100 g				
FRAGRANCE LAVENDER & CHIA F-153480 (UNII: SXS9CO2TZK)					0	0.02 g in 100 g				
р	ackaging									
1	uchuging									
#	Item Code	Package Description		Marketing Sta Date	rt Marketing End Date					
1	NDC:76693- 101-45	80 g in 1 PACKAGE; Type 5: Device Coated or Otherwise Combinities with Biologic			07/01/2020					
N	Iarketing I	nfor	mation							
Marketing Category		gory	Application Number or Monograph Citation Marke		ting Start Date	Marketing End Date				
N	unapproved drug other									

Labeler - Sourcery Ltd (800787645)

# Registrant - Sourcery Ltd (800787645)

Revised: 5/2020

Sourcery Ltd