

KLEENEX INSTANT HAND SANITIZER- alcohol solution
Kimberly-Clark Corporation

KLEENEX® INSTANT HAND SANITIZER

Drug Facts

Active Ingredient

Ethyl Alcohol 68% v/v

Purpose

Antiseptic

Use

Hand sanitizer to decrease bacteria on the skin.

Warnings

Flammable - Keep product away from fire or flame.

For External Use Only.

When using this product avoid contact with eyes; in case of contact, flush eyes with water.

Stop use & ask a doctor if irritation or redness develops and persists.

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.

Other Information

- Report serious side effects from this product to 1-877-561-6587.
- Do not store above 110°F (40°C).

Inactive Ingredients

Water, Carbomer, Fragrance, Aminomethyl Propanol, Glycerin

Questions?

1-888-346-4652

Distributed in the U.S. by
Kimberly-Clark
Global Sales, LLC, Roswell,
GA 30076-2199

PRINCIPAL DISPLAY PANEL - 236 mL Bottle Label

Kleenex®
BRAND
MARQUE

Instant Hand
Sanitizer

kills
germs†

8 fl oz (236 mL)

20-14-141-0-09

Kleenex[®]
BRAND
MARQUE

Instant Hand Sanitizer

kills
germs*

8 fl oz (236 mL)

20-14-141-0-08

Drug Facts

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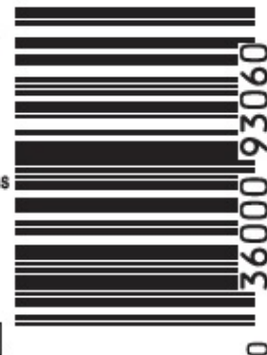
 **Kimberly-Clark**
PROFESSIONAL

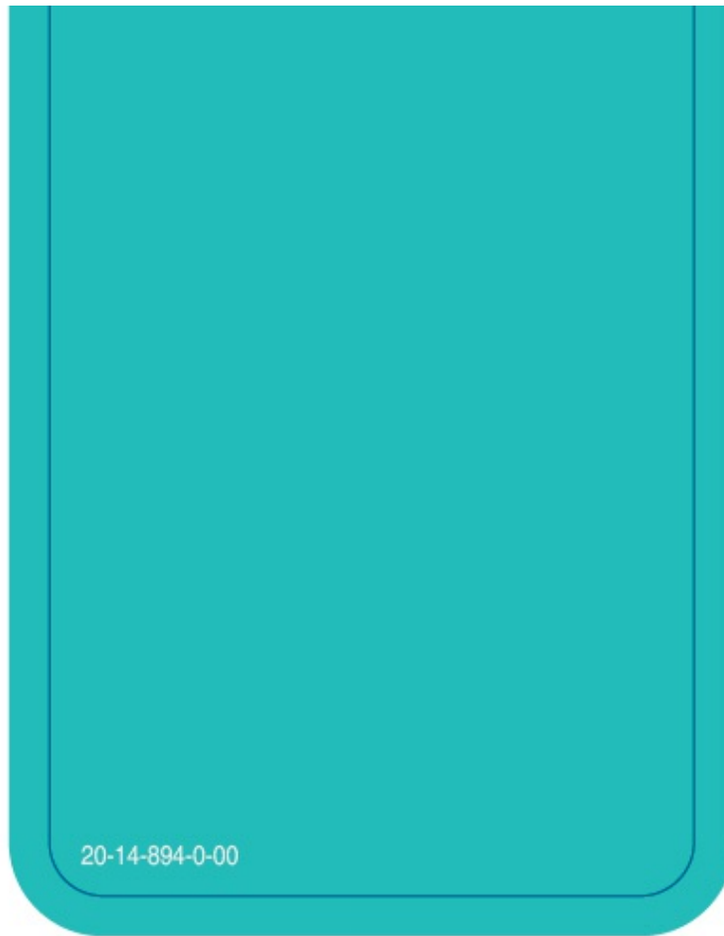
*Kills 99.99% of most common germs that make you sick.

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Made in Mexico of U.S. and/or Non-US materials / Fabriqué au Mexique à partir de matériaux américains et non américains
Distributed in the U.S. by Kimberly-Clark Global Sales, LLC, Roswell, GA 30076-2199
www.kcprofessional.com
Re-order #: 93060



NSF
Hardford Compounds
Program Listed E3
337322





KLEENEX INSTANT HAND SANITIZER

alcohol solution

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:55118-704
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
Alcohol (UNII: 3K9958V90M) (Alcohol - UNII:3K9958V90M)	Alcohol	68 mL in 100 mL

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
AMINOMETHYLPROPANOL (UNII: LU49E6626Q)	
GLYCERIN (UNII: PDC6A3C0OX)	
CARBOMER HOMOPOLYMER, UNSPECIFIED TYPE (UNII: 0A5MM307FC)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:55118-704-70	6 in 1 CARTON	08/30/2021	
1	NDC:55118-704-88	236 mL in 1 BOTTLE; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph drug	M003	08/30/2021	

Labeler - Kimberly-Clark Corporation (830997032)

Revised: 12/2024

Kimberly-Clark Corporation