ALOESEPT HS- ethanol liquid Walter G. Legge Company, Inc.

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.

Aloesept Hand Sanitizer

Drug Facts

Active Ingredient

Ethyl Alcohol 70% v/v

Purpose

Antiseptic

Uses

- For handwashing to decrease bacteria on the skin after changing diapers, assisting ill persons, or before contact with a person under medical care or treatment.
- Recommended for repeated use.

Warnings

• For external use only.

Keep out of eyes, ears, or mouth.

- Discontinue use if irritation occurs.
- Keep out of reach of children.

FLAMMABLE, KEEP AWAY FROM FIRE OR FLAME.

Directions

- If hands are visibly soiled, wash with soap and water and dry hands.
- Wet hands thoroughly with product, especially the area under the fingernails and allow to dry without rinsing.

Inactive ingredients benzalkonium chloride, chlorhexidine gluconate, isopropyl alcohol, PEG 10 dimethicone, PEG-14M, phenoxyethanol, polyquaternium 10, water.

Sold By: Walter G. Legge Company Inc.

444 Central Avenue, Peekskill, N.Y. 10566

1-800-345-3443

MADE IN THE USA

ALOESEPT HS

WATERLESS HAND SANITIZER

Compliant with CDC Hand Hygiene Guidelines.

Kills more than 99.99% of germs in 15 seconds.

Does not contain DEA/MEA, triclosan, parabens, formaldehyde, dyes or fragrances.

CONTAINS MOISTURIZING AGENTS

1000 ml



444 Central Ave., Peekskill, N.Y. 10588

www.leggesystems.com

Email: info@leggesystems.com

1-800-345-3443 Fax: 1-800-332-2838

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Questions? Call 1-800-345-3443 or visit our website www.leggesystems.com

5068-LEGGE-FL

Rev 101

ALOESEPT HS

1000 ml

MADE IN THE USA

ethanol liquid

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:63647-001

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name

Basis of Strength

ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)

ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)

70 mL in 100 mL

Ingredient Name Strength BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) CHLORHEXIDINE GLUCONATE (UNII: MOR84MUD8E) ISOPROPYL ALCOHOL (UNII: ND2M416302) PEG-10 DIMETHICONE (600 CST) (UNII: 8PR7V1SVM0) POLYETHYLENE OXIDE 600000 (UNII: 2126FD486L) PHENOXYETHANOL (UNII: HIE492ZZ3T)

POLYQUATERNIUM-10 (1000 MPA.S AT 2%) (UNII: GMR4PEN8PK)

WATER (UNII: 059QF0KO0R)

Packaging					
4	# Item Code	Package Description	Marketing Start Date	Marketing End Date	
]	NDC:63647- 001-18	532 mL in 1 BOTTLE; Type 0: Not a Combination Product	04/12/2021	07/13/2024	
2	NDC:63647- 001-34	1000 mL in 1 CARTRIDGE; Type 0: Not a Combination Product	04/12/2021	07/13/2024	

Marketing Information					
Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
part333E	04/12/2021	07/13/2024			
	Application Number or Monograph Citation	Application Number or Monograph Marketing Start Citation Date			

Labeler - Walter G. Legge Company, Inc. (001382274)

Revised: 9/2022 Walter G. Legge Company, Inc.