

OLIKA HYDRATING HAND SANITIZER FROSTED EUCALYPTUS- alcohol liquid
OLIKA HYDRATING HAND SANITIZER BLUE TANSY- alcohol liquid
OLIKA HYDRATING HAND SANITIZER COTTON FLOWER- alcohol liquid
Olika 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.

Olika Mindfulness Hand Sanitizers- 3 scents in 20 mL, 90 mL

This is a hand sanitizer manufactured according to the 1994 tentative final monograph for hand sanitizers and antiseptics. Final formulation finished product testing was performed to confirm efficacy and quality assurance.

The formulation comes in 2 package sizes: 20 mL and 90mL

Active Ingredient(s)

Ethyl Alcohol 65% v/v. Purpose: Antiseptic

Purpose

Antiseptic, Hand Sanitizer

Use

Helps reduce bacteria on skin.

Warnings

For external use only. Flammable. Keep away from heat or flame

Do not use

- in children less than 2 months of age
- on open skin wounds

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.

Directions

Spray liquid into hands and rub until dry. Use as often as needed.

Other information

- Store between 15-30C (59-86F)
- Avoid freezing and excessive heat above 40C (104F)

Inactive ingredients

aloe vera, fragrance, glycerin, hyaluronic acid, reishi mushrooms, water

20 mL Primary Container Spray Bottle Bottom Label

76751-711-02; 76751-712-02; 76751-713-02

20 mL Primary Container Spray Bottle Bottom Label



Package Label - Principal Display Panel and Information Panels

90 mL NDC: 76751-711-01

COOL 10

7655 U

2461 U

CMYK



ULTRA-HYDRATED HANDS

Many hand sanitizers dry out your hands, causing uncomfortable cracking or peeling that could expose you to more harmful bacteria. OLIKA's special blend of moisture-retaining hyaluronic acid and adaptogen-rich reishi quenches dry hands with long-lasting hydration while removing 99.9% of germs. Find out what's different about OLIKA.



Cruelty-free



More uses than similarly sized gels



Vegan formulation

This easy to use refill pouch will replenish your OLIKA three or four times for over 1500 sprays.

TO USE:

Simply unscrew the bottom of your OLIKA and fill with sanitizer up to the edge of the reservoir area.

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HAND SANITIZER

Learn more at [OLIKALIFE.COM](#)



EXP: 2024-JUN

Drug Facts	
Active Ingredient	Purpose
Ethyl Alcohol* 65% v/v	Antiseptic
Use Helps reduce bacteria on skin.	
Warnings	
For external use only. Flammable, keep away from fire or flame.	
Do not use ■ on children less than 12 months of age ■ on open skin wounds.	
When using keep out of eyes, ears, and mouth. In case of contact, 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.	
Other information	
Store between 15-30C (59-86F) ■ Avoid freezing and excessive heat above 40C (104F)	
Directions	
Refill empty bottle. ■ Spray liquid into hands and rub until dry. ■ Supervise children under 6 years of age to prevent swallowing.	
Inactive ingredients	
Aloe vera (aloe barbadensis) leaf juice, denatonium benzoate*, fragrance**, glycerin, hyaluronic acid, reishi mushroom, and water.	
Comments/Questions?	
(866) 446-5452 (866-44-OLIKA)	

*Alcohol denaturant. **for complete list visit [olikalife.com](#)

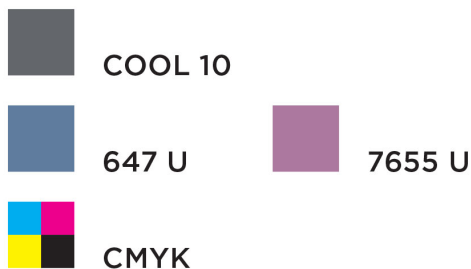
Package Label - Principal Display Panel and Information Panels

76751-711-02. 20 mL



Package Label - Principal Display Panel and Information Panels

76751-712-01 90 mL





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Designed in California Fragrance Made in USA Assembled in China Not tested on animals. NDC# 76751-712-01

HAND SANITIZER

Learn more at OLIKALIFE.COM



EXP: 2024-JUN

Drug Facts

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Directions

Refill empty bottle. ■ Spray liquid into hands and rub until dry. ■ Supervise children under 6 years of age to prevent swallowing.

Inactive ingredients

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Comments/Questions?

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*Alcohol denaturant. **For complete list visit olikalife.com

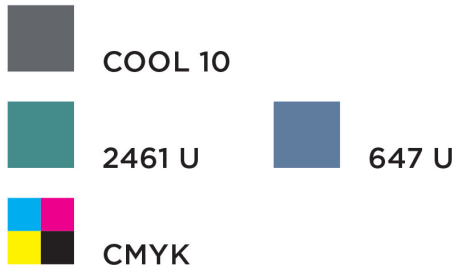
Package Label - Principal Display Panel and Information Panels

76751-712-02. 20 mL



Package Label - Principal Display Panel and Information Panels

76751-713-01. 90mL





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Designed in California
Fragrance Made in USA Not tested on animals.
Assembled in China NDC# 76751-713-01

HAND SANITIZER

Learn more at OLIKALIFE.COM



EXP. 2024-JUN

Drug Facts

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Ethyl Alcohol* 65% v/v	Antiseptic

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Directions

Refill empty bottles. ■ Spray liquid into hands and rub until dry. ■ Supervise children under 6 years of age to prevent swallowing.

Inactive ingredients

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Comments/Questions?

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*Alcohol denaturant. **for complete list visit olikalife.com

Package Label - Principal Display Panel and Information Panels

76751-713-02 20 mL



COOL 10 U



CMYK



2461 U



647 U (ALSO USE THIS COLOR TO FLOODCOAT THE INNER BOX)



OLIKA HYDRATING HAND SANITIZER FROSTED EUCALYPTUS

alcohol liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:76751-713
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	65 mL in 100 mL

Inactive Ingredients

Ingredient Name	Strength
ACETYL CEDRENE (UNII: X6I62755AK)	0.01667 mL in 100 mL
IONONE (UNII: QP734LIN1K)	0.00667 mL in 100 mL
ETHYL LINALOOL (UNII: SF2JS9GF5T)	0.005 mL in 100 mL
ETHYL BUTYRATE (UNII: UFD2LZ005D)	0.005 mL in 100 mL

FLORALOZONE (UNII: 1HA71K8K9L)	0.00417 mL in 100 mL
METHYL BENZODIOXEPINONE (UNII: 0NQ136C313)	0.00167 mL in 100 mL
GLYCERIN (UNII: PDC6A3C0OX)	4.2 mL in 100 mL
WATER (UNII: 059QF0KOOR)	29.78 mL in 100 mL
NERYL ACETATE (UNII: OF82IJU18H)	0.00833 mL in 100 mL
DIPROPYLENE GLYCOL (UNII: E107L85C40)	0.14825 mL in 100 mL
2-ISOBUTYL-4-METHYLTETRAHYDROPYRAN-4-OL (UNII: VK5ZHH2T3F)	0.08333 mL in 100 mL
ALLYL HEPTANOATE (UNII: AU4CYG9V68)	0.00033 mL in 100 mL
METHYL DIHYDROJASMONATE (SYNTHETIC) (UNII: 3GW44CIE3Y)	0.075 mL in 100 mL
DIHYDROMYRCENOL (UNII: 46L1B02ND9)	0.03333 mL in 100 mL
ETHYLENE BRASSYLATE (UNII: 9A87HC7ROD)	0.06667 mL in 100 mL
3-ISOCAMPHYL CYCLOHEXANOL, TRANS- (UNII: 34UP96K73Z)	0.025 mL in 100 mL
HEXYL ACETATE (UNII: 7U7KU3MWT0)	0.00667 mL in 100 mL
GERANYL ACETATE (UNII: 3W81YG7P9R)	0.005 mL in 100 mL
ETHYL ACETOACETATE ETHYLENEGLYCOL KETAL (UNII: G5EXI4NID0)	0.00167 mL in 100 mL
PENTADECALACTONE (UNII: OK17S3S98K)	0.00167 mL in 100 mL
CYCLOPENTANONE (UNII: 220W81TN3S)	0.00167 mL in 100 mL
ETHYL 2,4-DECADIENOATE, (2E,4Z)- (UNII: 79P6KS9Y5Z)	0.00083 mL in 100 mL
4-(P-HYDROXYPHENYL)-2-BUTANONE (UNII: 7QY1MH15BG)	0.00083 mL in 100 mL
HYALURONATE SODIUM (UNII: YSE9PPT4TH)	0.07 mL in 100 mL
ALLYL .ALPHA.-IONONE (UNII: 8IP66F9ODG)	0.00083 mL in 100 mL
ALLYL HEXANOATE (UNII: 3VH84A363D)	0.00083 mL in 100 mL
ALLYL CYCLOHEXANEACETATE (UNII: M6J8835739)	0.00058 mL in 100 mL
3-HEXEN-1-OL, (3Z)- (UNII: V14F8G75P4)	0.0005 mL in 100 mL
ALOE VERA LEAF (UNII: ZY81Z83H0X)	0.25 mL in 100 mL
REISHI (UNII: TKD8LH0X2Z)	0.2 mL in 100 mL

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:76751-713-02	1 in 1 BLISTER PACK	06/14/2021	
1	NDC:76751-713-00	20 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product		
2	NDC:76751-713-01	90 mL in 1 POUCH; Type 0: Not a Combination Product	06/14/2021	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	06/14/2021	

OLIKA HYDRATING HAND SANITIZER BLUE TANSY

alcohol liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:76751-712
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	65 mL in 100 mL

Inactive Ingredients

Ingredient Name	Strength
ETHYLENE BRASSYLATE (UNII: 9A87HC7ROD)	0.1021 mL in 100 mL
TETRAMETHYL ACETYLOCTAHYDRONAPHTHALENES (UNII: 2JU6ZH6GRE)	0.0592 mL in 100 mL
DIPROPYLENE GLYCOL (UNII: E107L85C40)	0.1567 mL in 100 mL
ETHYL 2-METHYLBUTYRATE (UNII: L1T4AB29DS)	0.00002 mL in 100 mL
METHYL DIHYDROJASMONATE (SYNTHETIC) (UNII: 3GW44CIE3Y)	0.0828 mL in 100 mL
2-ISOBUTYL-4-METHYLTETRAHYDROPYRAN-4-OL (UNII: VK5ZHH2T3F)	0.0237 mL in 100 mL
CYCLODODECANE (UNII: 97CN13ZD83)	0.0207 mL in 100 mL
2-TERT-BUTYLCYCLOHEXYL ACETATE (UNII: 364FV60913)	0.008 mL in 100 mL
NAPHTHO(2,1-B)FURAN (UNII: 5O098D6W4T)	0.0033 mL in 100 mL
ALOE VERA LEAF (UNII: ZY81Z83H0X)	0.25 mL in 100 mL
GLYCERIN (UNII: PDC6A3C0OX)	4.2 mL in 100 mL
WATER (UNII: 059QF0KO0R)	29.78 mL in 100 mL
4-HYDROXY-1-(5-ISOCAMPHYL)CYCLOHEXANE (UNII: TVJ25725XP)	0.0142 mL in 100 mL
DIETHYL MALONATE (UNII: 53A58PA183)	0.0104 mL in 100 mL
2,4-DIMETHYL-4-NONANOL (UNII: 625246Q74Q)	0.0053 mL in 100 mL
4-TERT-BUTYLCYCLOHEXYL ACETATE (UNII: 21EUM2B8UC)	0.0044 mL in 100 mL
3-ISOCAMPHYLCYCLOHEXANOL, TRANS- (UNII: 34UP96K73Z)	0.0036 mL in 100 mL
CEDRYL ACETATE (UNII: 0WS0WJ9WNV)	0.0015 mL in 100 mL
AMBROXIDE, (-)- (UNII: TD34B3O8M9)	0.0009 mL in 100 mL
METHYL 3-METHYLORSELLINATE (UNII: 12YH9T04QE)	0.0006 mL in 100 mL
4-METHYL-3-DECEN-5-OL, (3E)- (UNII: 5S5I61TY7P)	0.0004 mL in 100 mL
DIHYDRODEHYDRO-.BETA.-IONONE (UNII: XZS4VD987O)	0.0002 mL in 100 mL

ETHYL 2-METHYLPENTANOATE (UNII: 405SN8638D)	0.0002 mL in 100 mL
.GAMMA.-DECALACTONE (UNII: 7HLS05KP9O)	0.0001 mL in 100 mL
DECAHYDRO-2,2,6,6,7,8,8-HEPTAMETHYL-2H-INDENO(4,5-B)FURAN (UNII: 89ZYZ7YI66)	0.00004 mL in 100 mL
METHYL ANTHRANILATE (UNII: 981I0C1E5W)	0.00002 mL in 100 mL
2,4-DIMETHYL-3-CYCLOHEXENE CARBOXALDEHYDE (UNII: 452GFV2AFS)	0.0002 mL in 100 mL
3-HEXEN-1-OL, (3Z)- (UNII: V14F8G75P4)	0.0004 mL in 100 mL
HEX-3-EN-1-YL METHYL CARBONATE, (3Z)- (UNII: WEC5YY1PGT)	0.0004 mL in 100 mL
DIMETHYLCYCLOHEXYLETHOXY ISOBUTYLPROPANOATE (UNII: I1AI9W9CAB)	0.0003 mL in 100 mL
ETHYL MALTOL (UNII: L6Q8K29L05)	0.0003 mL in 100 mL
ALLYL CYCLOHEXANEACETATE (UNII: M6J8835739)	0.00003 mL in 100 mL
1-METHYL-2-(((1R,3S,5S)-1,2,2-TRIMETHYLBICYCLO HEX-3-YL)METHYL)CYCLOPROPANEMETHANOL, (1R,2R)- (UNII: 56FP2ES9B7)	0.00003 mL in 100 mL
REISHI (UNII: TKD8LH0X2Z)	0.2 mL in 100 mL
HYALURONATE SODIUM (UNII: YSE9PPT4TH)	0.07 mL in 100 mL

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:76751-712-02	1 in 1 BLISTER PACK	06/14/2021	
1	NDC:76751-712-20	20 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product		
2	NDC:76751-712-01	90 mL in 1 POUCH; Type 0: Not a Combination Product	06/14/2021	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	06/14/2021	

OLIKA HYDRATING HAND SANITIZER COTTON FLOWER

alcohol liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:76751-711
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	65 mL in 100 mL

Inactive Ingredients

Ingredient Name	Strength
METHYL DIHYDROJASMONATE (SYNTHETIC) (UNII: 3GW44CIE3Y)	0.0548 mL in 100 mL
2,2,5-TRIMETHYLHEXANE (UNII: AGW1Z6985E)	0.0103 mL in 100 mL
DIHYDRODEHYDRO-.BETA.-IONONE (UNII: XZS4VD987O)	0.0008 mL in 100 mL
2,4-DIMETHYL-3-CYCLOHEXENE CARBOXALDEHYDE (UNII: 452GFV2AFS)	0.0007 mL in 100 mL
.GAMMA.-DECALACTONE (UNII: 7HLS05KP9O)	0.0007 mL in 100 mL
.ALPHA.-METHYLBENZYL ACETATE (UNII: FYS3E9NBA3)	0.0007 mL in 100 mL
ETHYL 2,2-DIMETHYLHYDROCINNAMAL (UNII: 5V2FN5AA3W)	0.0003 mL in 100 mL
2-ISOBUTYL-4-METHYLTETRAHYDROPYRAN-4-OL (UNII: VK5ZHH2T3F)	0.0685 mL in 100 mL
4-HYDROXY-1-(5-ISOCAMPHYL)CYCLOHEXANE (UNII: TVJ25725XP)	0.0034 mL in 100 mL
.GAMMA.-UNDECALACTONE (UNII: QB1T0AG2YL)	0.0007 mL in 100 mL
NAPHTHO(2,1-B)FURAN (UNII: 5O098D6W4T)	0.0007 mL in 100 mL
3-HEXENYL ACETATE, (3Z)- (UNII: 6INA6GC5I6)	0.0003 mL in 100 mL
DIPROPYLENE GLYCOL (UNII: E107L85C40)	0.1603 mL in 100 mL
TETRAMETHYL ACETYLOCTAHYDRONAPHTHALENES (UNII: 2JU6ZH6GRE)	0.0836 mL in 100 mL
ETHYLENE BRASSYLATE (UNII: 9A87HC7ROD)	0.0753 mL in 100 mL
PHENYLETHYL ALCOHOL (UNII: ML9LGA7468)	0.0075 mL in 100 mL
IONONE (UNII: QP734LIN1K)	0.0034 mL in 100 mL
CYCLOHEXANEPROPANOL, 2,2,6-TRIMETHYL-.ALPHA.-PROPYL- (UNII: CLV4EM4325)	0.0021 mL in 100 mL
2-TERT-BUTYLCYCLOHEXYL ACETATE (UNII: 364FV60913)	0.001 mL in 100 mL
ALLYL HEPTANOATE (UNII: AU4CYG9V68)	0.0005 mL in 100 mL
REISHI (UNII: TKD8LH0X2Z)	0.2 mL in 100 mL
HYALURONATE SODIUM (UNII: YSE9PPT4TH)	0.07 mL in 100 mL
ALOE VERA LEAF (UNII: ZY81Z83H0X)	0.25 mL in 100 mL
DIHYDROMYRCENOL (UNII: 46L1B02ND9)	0.0199 mL in 100 mL
2-TERT-BUTYLCYCLOHEXYLOXYBUTANOL (UNII: 1DR20642YH)	0.0034 mL in 100 mL
JASMONE (UNII: RC4W0G9YUK)	0.0006 mL in 100 mL
CYCLOPENTANONE (UNII: 220W81TN3S)	0.0003 mL in 100 mL
TRIMETHYLBENZYLSILANE (UNII: HFX8B6Q8WC)	0.0002 mL in 100 mL
GLYCERIN (UNII: PDC6A3C0OX)	4.2 mL in 100 mL
WATER (UNII: 059QF0KO0R)	29.78 mL in 100 mL

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:76751-711-02	1 in 1 BLISTER PACK	06/14/2021	
1	NDC:76751-711-20	20 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product		
2	NDC:76751-711-01	90 mL in 1 POUCH; Type 0: Not a Combination Product	06/14/2021	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	06/14/2021	

Labeler - Oliko Inc. (080476192)

Revised: 12/2021

Oliko Inc.