# PURELL ADVANCED E3 RATED INSTANT HAND SANITIZER- alcohol liquid GOJO Industries, 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.

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#### PURELL Advanced E3 Rated Instant Hand Sanitizer

#### Active ingredient

Ethyl Alcohol 70% v/v

#### Purpose

Antimicrobial

#### Uses

- Hand sanitizer to help reduce bacteria on the skin that could cause disease
- Recommended for repeated use

#### Warnings

#### Flammable. Keep away from fire or flame.

#### For external use only

**When using this product** do not use in or near the eyes. In case of contact, rinse eyes thoroughly with water.

Stop use and ask a doctor if irritation or rash appears and lasts

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

#### Directions

- Place enough product in your palm to thoroughly cover your hands
- Rub hands together briskly until dry
- No rinsing required
- No towels needed

#### **Inactive ingredients**

Water (Aqua), Isopropyl Alcohol, Caprylyl Glycol, Glycerin, Isopropyl Myristate, Tocopheryl Acetate, Acrylates/C10-30 Alkyl Acrylate Crosspolymer, Aminomethyl Propanol



#### PURELL ADVANCED E3 RATED INSTANT HAND SANITIZER

alcohol liquid

| Product Information             |  |         |                       |               |  |  |
|---------------------------------|--|---------|-----------------------|---------------|--|--|
| Product T ype                   | HUMAN OTC DRUG Item Code (Source) NDC: |         |                       | NDC:21749-706 |  |  |
| Route of Administration         | TOPICAL                                |         |                       |               |  |  |
|                                 |  |         |                       |               |  |  |
| Active Ingredient/Active Moiety |  |         |                       |               |  |  |
| Ingree                          | Basis of Strength                      |         | h Strength            |               |  |  |
| ALCOHOL (UNII: 3K9958V90M) (ALC |  | ALCOHOL | $0.70\ mL$ in $1\ mL$ |               |  |  |
|                                 |  |         |                       |               |  |  |
| Inactive Ingredients            |  |         |                       |               |  |  |
|                                 | Strength                               |         |                       |               |  |  |

| Water (UNII: 059QF0K   | ,   |                      |                    |  |  |  |  |
|--|---|----------------------|--------------------|--|--|--|--|
|  | Isopropyl Alcohol (UNII: ND2M416302)  |                      |                    |  |  |  |  |
| Glycerin (UNII: PDC6A3C0OX)  |   |                      |                    |  |  |  |  |
| Isopropyl Myristate (U   | Isopropyl Myristate (UNII: 0RE8K4LNJS)  |                      |                    |  |  |  |  |
| .ALPHATO COPHEROL ACETATE, D- (UNII: A7E6112E4N)                   |   |                      |                    |  |  |  |  |
| Caprylyl Glycol (UNII: 00 YIU5438 U)                               |   |                      |                    |  |  |  |  |
| AMINO METHYLPRO  | PANOL (UNII: LU49E6626Q)  |                      |                    |  |  |  |  |
|  |   |                      |                    |  |  |  |  |
|  |   |                      |                    |  |  |  |  |
| Packaging  |   |                      |                    |  |  |  |  |
| # Item Code  | Package Description   | Marketing Start Date | Marketing End Date |  |  |  |  |
| 1 NDC:21749-706-08   | 237 mL in 1 BOTTLE; Type 0: Not a Combination Product   | 0 3/31/20 13         |                    |  |  |  |  |
| <b>2</b> NDC:21749-706-12  | 355 mL in 1 BOTTLE; Type 0: Not a Combination Product   | 0 3/31/20 13         |                    |  |  |  |  |
| 3 NDC:21749-706-97   | 700 mL in 1 BOTTLE; Type 0: Not a Combination Product   | 03/31/2013           |                    |  |  |  |  |
| 5 1120.21/45 /00 5/  | , oo mii m i bo i iii, iype o. Not a Combination i fouaet   | 03/31/2013           |                    |  |  |  |  |
|  | 1000 mL in 1 BOTTLE; Type 0: Not a Combination Product  |                      |                    |  |  |  |  |
| 4 NDC:21749-706-10   |   | 03/31/2013           |                    |  |  |  |  |
| <b>4</b> NDC:21749-706-10  | 1000 mL in 1 BOTTLE; Type 0: Not a Combination Product  | 03/31/2013           |                    |  |  |  |  |
| 4 NDC:21749-706-10   | 1000 mL in 1 BOTTLE; Type 0: Not a Combination Product  | 03/31/2013           |                    |  |  |  |  |
| 4 NDC:21749-706-10   | 1000 mL in 1 BOTTLE; Type 0: Not a Combination Product<br>1200 mL in 1 BOTTLE; Type 0: Not a Combination Product                    | 03/31/2013           |                    |  |  |  |  |
| <ul> <li>4 NDC:21749-706-10</li> <li>5 NDC:21749-706-89</li> </ul> | 1000 mL in 1 BOTTLE; Type 0: Not a Combination Product<br>1200 mL in 1 BOTTLE; Type 0: Not a Combination Product<br><b>Drmation</b> | 03/31/2013           | Marketing End Date |  |  |  |  |

## Labeler - GOJO Industries, Inc. (004162038)

# EstablishmentNameAddressID/FEIBusiness OperationsGOJO Industries, Inc.036424534manufacture(21749-706)

### Establishment

| Name                  | Address | ID/FEI    | Business Operations                                       |
|-----------------------|---------|-----------|---|
| GOJO Industries, Inc. |         | 088312414 | manufacture(21749-706), label(21749-706), pack(21749-706) |

Revised: 12/2019

GOJO Industries, Inc.