

**PRO ADVANTAGE INSTANT HAND SANITIZER - alcohol liquid**  
**NDC National Distribution & Contracting, 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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**Pro Advantage Instant Hand Sanitizer**

**Drug Facts**

**ACTIVE INGREDIENT:**

Alcohol, 62%

**Purpose**

Antiseptic

**USE**

To help reduce bacteria on the hands that can potentially cause disease.

**WARNING:**

- For External Use Only
- Flammable, Keep away from fire or flame.

**When using this product**

- Avoid contact with eyes, if this occurs rinse thoroughly with water and contact a physician.

**Ask a doctor before use if you have**

- deep wounds, animal bites or serious burns.

**Stop use and ask a doctor if**

- condition persists.

**Keep out of reach of children.**

- If swallowed get medical help or contact a Poison Control Center right away.

**DIRECTIONS:**

Apply a liberal amount to hands and rub hands thoroughly until dry. Do not rinse or wipe off gel.

**INACTIVE INGREDIENTS**

Water, Glycerin, Fragrance, Carbomer, Triethanolamine, DMDM Hydantoin

REF: P779114 NDC 43128-114-03

Made in China  
www.ProAdvantagebyNDC.com

Manufactured for NDC, Inc.  
407 New Sanford Road, La Vergne, TN 37086

Pro Advantage by NDC Instant  
Hand Sanitizer  
4 FL. OZ.  
( 118 mL )  
Kills Germs With 62% Alcohol  
\*+M220P77911480\*

**Product Labels**

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alcohol liquid

## Product Information

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:43128-114
<b>Route of Administration</b>	TOPICAL		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	620 mg in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
GLYCERIN (UNII: PDC6A3C0OX)	
TROLAMINE (UNII: 9O3K93S3TK)	
DMDM HYDANTOIN (UNII: BYR0546TOW)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:43128-114-03	1 in 1 PACKAGE		
1		118 mL in 1 BOTTLE		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333E	03/08/2013	

**Labeler** - NDC National Distribution & Contracting, Inc. (009831413)

Revised: 3/2013

NDC National Distribution & Contracting, Inc.