# WITCH HAZEL- witch hazel liquid Pharmacy Vlaue Alliance LLC

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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Witch Hazel 822.001/822AA

### **Active ingredient**

Witch hazel 86%

#### **Purpose**

Astringent

#### use

for relief of minor skin irritations due to:

- insects bites
- minor cuts
- minor scrapes

### Warnings

For external use only

### When using this product

avoid contact with the eyes

### Stop use and ask a doctor if

• condition worsens or symptoms persist for more than 7 days

### Keep out of reach of children

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

#### **Directions**

apply as often as needed

### inactive ingredients

#### Questions

Call 1-888-593-0593

\*This product is not manufactued or distributed by Dickinson Brand, Inc., distributorof T.N. Dickinson Witch Hazel.

If for any reason you are not satisfied with this product, please return it to the store where purchased for a full refund.

Distributed by: Phamacy Value Alliance LLC

407 East Lancaster Avenue, Wayne, PA 19087

www.emersongroup.com

### **Principal Panel Display**

COMPARE TO T.N.DICKINSON'S\*

Premier Value Witch Hazel u.s.p.

astringent

hamamelis water

for relief of minor skin irritations due to:

- insects bites
- minor cuts
- minor scrapes

Square bottle uses less plastic than a similarly sized round bottle

Recyclable (if available in your area)

16 FL OZ (1 PT) 473 mL



### **WITCH HAZEL**

witch hazel liquid

#### **Product Information**

Product Type HUMAN OTC DRUG Item Code (Source) NDC:68016-822

Route of Administration TOPICAL

#### **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
WITCH HAZEL (UNII: 1011410U34) (WTCH HAZEL - UNII:1011410U34)	WTCH HAZEL	842 mg in 1 ml

Inactive Ingredients			
Ingredient Name	Strength		
ALCOHOL (UNII: 3K9958V90M)			

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
	NDC:68016- 822-43	473 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	08/15/2003		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part347	08/15/2003	

## Labeler - Pharmacy Vlaue Alliance LLC (101668460)

# Registrant - Vi-Jon, LLC (790752542)

Establishment				
Name	Address	ID/FEI	<b>Business Operations</b>	
Vi-Jon, LLC		790752542	manufacture(68016-822)	

Establishment				
Name	Address	ID/FEI	<b>Business Operations</b>	
Vi-Jon, LLC		088520668	manufacture(68016-822)	

Revised: 9/2022 Pharmacy Vlaue Alliance LLC