HYDROCORTISONE- hydrocortisone lotion AKRON PHARMA 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.

Hydrocortisone USP 1% Lotion

Drug Facts

Active Ingredient

Hydrocortisone 1% (Micronized)

Purpose

Antipruritic (Anti-itch)

Use

For the temporary relief of minor skin irritations, inflammations, itching and rashes caused by:

- insect bites
- eczema
- psoriasis
- soaps
- detergents
- cosmetics,
- jewelry,
- poison oak,
- poison sumac
- Other uses of this product should be undertaken only under the advice and supervision of a doctor.

Warnings

For external use only.

When using this product

Do not get into eyes. If contact occurs, rinse thoroughly with water.

Stop use and ask a doctor if

- Condition worsens
- If symptoms persist for more than 7 days or clear up and occur again within a few days. Discontinue use of this product and do not begin use of any other hydrocortisone product unless you have consulted a doctor.
- Do no use for diaper rash. Consult a doctor.

Keep out of Reach of Children

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

Directions

- Shake will before using.
- For adults and children 2 years of age and older: Apply to affected area not more than 2 to 4 times daily.
- For children under 2 years of age: there is no recommended dosage except under the advice and supervision of a doctor.

Other Information

• Store away from excessive heat or cold. Shake well before using.

Inactive Ingredients

Purified Water, Glycerin, Cetyl Alcohol, Benzyl Alcohol, Sodium Lauryl Sulfate, Stearyl Alcohol, Aloevera, Xanthan Gum

Questions?

Please Call 1(877) 225-6999

Manufactured for:

Akron Pharma, Inc. Fairfield, NJ 07004

Rev. 03/19

Manufactured in U.S.A

NDC 71399-0120-1		Drug Facts Active Ingredient Purpose	
Hudrocorticono	USD 10/	Active Ingredient Purpose Hydrocortisone USP 1% (Micronized) Antipruritic (Anti-itch) Use Use For the temporary relief of minor skin irritations, inflammations, itches and rashes caused by:=seborrheic dermatitis = insect bites = eczema = psoriasis = soaps edetergents = cosmetics = jewelry = poison oak poison ivy = poison sumac = other uses of this product should be undertaken only under the advice and supervision of a doctor.	
Lotion		Warnings For external use only. When using this product Do not get into eyes. If contact occurs, rinse thoroughly with water.	
With Aloe Antipruritie	Micronized c (Anti-Itch)	Stop use and ask a doctor if Condition worsens = If symptoms persist for more than 7 days or clear up and occur again within a few days, discontinue use of this product and do not begin use of any other hydrocortisone product unless you have consulted a doctor. = Do no use for diaper rash. Consult a doctor.	
		 Keep out of Reach of Children If swallowed, get medical help or contact Poison Control Center right away. Directions - Shake well before using For adults and children 2 years of age and older: Apply to affected area not more than 3 to 4 times daily For children under 2 years of age: there is no recommended dosage except under the advice and supervision of a doctor. 	
For external use only.		Other Information - Store away from excessive heat or cold. Shake well before using.	
		Inactive Ingredients Purified Water, Glycerin, Cetyl Alcohol, Benzyl Alcohol, Sodium Lauryl Sulfate, Stearyl Alcohol, Aloe Vera, Xanthan Gum.	
		Questions? Please call 1 (877) 225-6999	
4 fl. oz. (120ml)	Akrðn Pharma	Manufactured In U.S.A Manufactured for: Akron Pharma, Inc., Fairfield, NJ 07004 Rev. 03/19 7 13 990 12 00 1	

HYDROCORTISONE

hydrocortisone lotion **Product Information** HUMAN OTC DRUG NDC:71399-0120 **Product Type** Item Code (Source) TOPICAL **Route of Administration Active Ingredient/Active Moiety** Ingredient Name **Basis of Strength** Strength HYDROCORTISONE (UNII: W4X0X7BPJ) (HYDROCORTISONE - UNII:W4X0X7BPJ) HYDROCORTISONE 10 mg in 1 mL **Inactive Ingredients** Strength **Ingredient Name** WATER (UNII: 059QF0KO0R) GLYCERIN (UNII: PDC6A3C0OX) CETYL ALCOHOL (UNII: 936JST6JCN)

SC	DDIUM LAURYL S					
ST	EARYL ALCOHOI	_ (UNII: 2KR89I4H1Y)				
AL	OE VERA LEAF (U					
XA	ANTHAN GUM (UN					
Packaging						
#	ltem Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:71399- 0120-1	1 in 1 CARTON	05/17/2021			
1		120 mL in 1 BOTTLE; Type 0: Not a Combination Product				
Marketing Information						
	Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
	FC monograph not		05/17/2021			

Labeler - AKRON PHARMA INC (067878881)

Revised: 3/2023

AKRON PHARMA INC