AQUANIL HC- hydrocortisone lotion Person and Covey

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.

Aquanil HC

Active Ingredient

Hydrocortisone

Stop Use and Ask a Doctor

Stop use and ask a doctor if:

- -Condition worsens
- -If symptoms persit for more than 7 days or clear up and occur again within a few days. Discocntinue 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 the Reach of Children

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

Purpose

Antipruritic (Anti-itch)

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 ddoctor.
- -Store away from excessive heat or cold.

Inactive Ingredients

Purified Water, Glycerin, Cetyl Alcohol, Benzyl Alcohol, Sodium Laureth Sulfate, Stearyl

Questions?

Questions? Please call (800) 423-2341

Uses

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

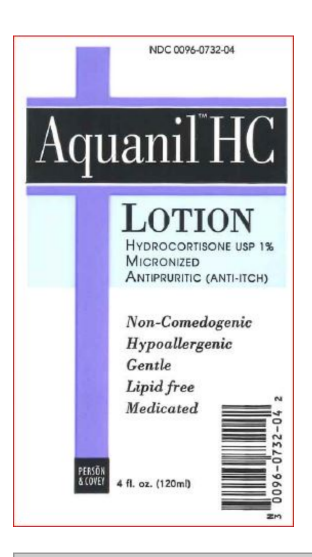
- -insect bites
- -eczema
- -psoriasis
- -soaps
- -detergents
- -cosmetics,
- -jewelry,
- -poison oak,
- -poison sumac
- -Other uses of this product should be undertaken only under the advice and supervision of a doctor.

Warnings

For external use only.

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

Package Label. Principal Display Panel



AQUANIL HC

hydrocortisone lotion

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Product Type HUMAN OTC DRUG Item Code (Source) NDC:0096-0732

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name Basis of Strength Strength

HYDROCORTISONE (UNII: W4X0X7BPJ) (HYDROCORTISONE - UNII:W4X0X7BPJ) HYDROCORTISONE 0.01 g in 1 g

Inactive Ingredients

Ingredient Name	Strength
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BENZYL ALCOHOL (UNII: LKG8494WBH)

WATER (UNII: 059QF0KO0R)

SODIUM LAURYL SULFATE (UNII: 368GB5141J)

XANTHAN GUM (UNII: TTV12P4NEE)
GLYCERIN (UNII: PDC6A3C0OX)
CETYL ALCOHOL (UNII: 936JST6JCN)

CTEADVI	ALCOHOL	/LINIII.	2KR89I4H1Y)
SIEARTL	ALCURUL	(UIMII:	/KK6914H111

Packaging				
	# Item Code Package Description		Marketing Start Date	Marketing End Date
	NDC:0096-0732-	118 g in 1 BOTTLE; Type 0: Not a Combination Product	01/08/1995	
	NDC:0096-0732- 15	16 g in 1 BOTTLE; Type 0: Not a Combination Product	01/08/1995	01/07/2022

Marketing Information			
Marketing Application Number or Monograph Category Citation		Marketing Start Date	Marketing End Date
OTC monograph not final	part348	01/08/1995	
	part340	01/00/1993	

Labeler - Person and Covey (008482473)

Establishment				
Na me	Address	ID/FEI	Business Operations	
Person and Covey		008482473	manufacture(0096-0732)	

Revised: 10/2022 Person and Covey