DRS. HYDROCORTISONE ALOE VERA- hydrocortisone cream OL PHARMA TECH,LLC

ACTIVE INGREDIENT

Hydrocortisone, USP 1%

PURPOSE

Anti-itch

USES

for the temporary relief of itching associated with minor skin irritations, inflammation, and rashes due to:

- eczema
- insect bites
- poison ivy
- •poison oak
- •poison ...

WARNINGS

For external use only

DO NOT USE

for external feminine itching if you have a vaginal discharge. Consult a doctor. for the treatment of diaper rash. Consult a doctor.

WHEN USING THIS PRODUCT

avoid contact with the eyes

•do not begin the use of any other hydrocortisone product unless directed by a doctor

•for external anal itching:

do not use more than directed unless directed by a doctor

do not put this product into the rectum by using fingers or any mechanical device or applicator

STOP USE AND ASK YOUR DOCTOR IF

- •symptoms last for more than 7 days
- •the condition gets worse
- •symptoms clear up and occur again in a few days
- •rectal bleeding occurs, consult doctor promptly

KEEP OUT OF REACH OF CHILDREN

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

DIRECTIONS

- Adults and children 2 years of age and older: apply to affected area not more than 3 to 4 times daily
- Children under 2 years of age: do not use, consult a doctor
- For external anal itching
- Adults: when practical, clean the affected area with mild soap and warm water, rinse thoroughly, gently dry by patting or blotting with toilet tissue or a soft cloth before application of this product

OTHER INFORMATION

- Store at room temperature 59°-86°F (15°-30°C).
- Protect from freezing.
- Before using any medication, read all label directions. Keep carton, it contains important information.

INACTIVE INGREDIENTS

cetostearyl alcohol, sodium cetostearyl sulfate, stearic acid, trolamine, mineral oil , propylene glycol, water, methyl paraben, propyl paraben, EDTA, vitamin E, Aloe vera leaf.



hydrocortisone cream					
Product Information					
Product Type	HUMAN OTC DRUG	em Code (So	urce)	NDC:80	489-088
Route of Administration	TOPICAL				
Active Ingredient/Active	Moiety				
Ingredient Name			Basis of Str	rength	Strength
HYDROCORTISONE (UNII: W4X0	(7BPJ) (HYDROCORTISONE - UNI	II:WI4X0X7BPJ)	HYDROCORTIS	ONE	10 mg in 1 g
Inactive Ingredients					
Inactive Ingredients					
Inactive Ingredients	Ingredient Name			S	trength
Inactive Ingredients CETEARETH-12 (UNII: 7V4MR24V	-			S	trength
CETEARETH-12 (UNII: 7V4MR24V PROPYLENE GLYCOL (UNII: 6DC	5P) 9Q167V3)			S	trength
CETEARETH-12 (UNII: 7V4MR24V PROPYLENE GLYCOL (UNII: 6DC METHYLPARABEN (UNII: A218C7F	5P) 9Q167V3) II9T)			S	trength
CETEARETH-12 (UNII: 7V4MR24V PROPYLENE GLYCOL (UNII: 6DC METHYLPARABEN (UNII: A218C7F EDETATE DISODIUM (UNII: 7FLD	5P) 9Q167V3) II9T)			S	trength
CETEARETH-12 (UNII: 7V4MR24V PROPYLENE GLYCOL (UNII: 6DC METHYLPARABEN (UNII: A218C7F EDETATE DISODIUM (UNII: 7FLD TROLAMINE (UNII: 903K93S3TK)	5P) 9Q167V3) II9T) 91C86K)			S	trength
CETEARETH-12 (UNII: 7V4MR24V PROPYLENE GLYCOL (UNII: 6DC METHYLPARABEN (UNII: A2I8C7F EDETATE DISODIUM (UNII: 7FLD TROLAMINE (UNII: 903K93S3TK) SODIUM CETOSTEARYL SULFA	5P) 9Q167V3) II9T) 91C86K) FE (UNII: 7ZBS06BH4B)			S	trength
CETEARETH-12 (UNII: 7V4MR24V PROPYLENE GLYCOL (UNII: 6DC METHYLPARABEN (UNII: A218C7F EDETATE DISODIUM (UNII: 7FLD TROLAMINE (UNII: 903K93S3TK) SODIUM CETOSTEARYL SULFA PROPYLPARABEN (UNII: Z8IX2SC	5P) 9Q167V3) II9T) 91C86K) FE (UNII: 7ZBS06BH4B)			S	trength
CETEARETH-12 (UNII: 7V4MR24V PROPYLENE GLYCOL (UNII: 6DC METHYLPARABEN (UNII: A2I8C7F EDETATE DISODIUM (UNII: 7FLD TROLAMINE (UNII: 903K93S3TK) SODIUM CETOSTEARYL SULFA PROPYLPARABEN (UNII: Z8IX2SC WATER (UNII: 059QF0K00R)	5P) 9Q167V3) II9T) 91C86K) FE (UNII: 7ZBS06BH4B) :1OH)			S	trength
CETEARETH-12 (UNII: 7V4MR24V PROPYLENE GLYCOL (UNII: 6DC METHYLPARABEN (UNII: A218C7F EDETATE DISODIUM (UNII: 7FLD TROLAMINE (UNII: 903K93S3TK) SODIUM CETOSTEARYL SULFA PROPYLPARABEN (UNII: Z8IX2SC WATER (UNII: 059QF0K00R) CETOSTEARYL ALCOHOL (UNII:	5P) 9Q167V3) II9T) 91C86K) FE (UNII: 7ZBS06BH4B) 10H) 2DMT128M1S)			S	trength
CETEARETH-12 (UNII: 7V4MR24V PROPYLENE GLYCOL (UNII: 6DC METHYLPARABEN (UNII: A218C7F EDETATE DISODIUM (UNII: 7FLD TROLAMINE (UNII: 903K93S3TK) SODIUM CETOSTEARYL SULFA PROPYLPARABEN (UNII: Z81X2SC WATER (UNII: 059QF0K00R) CETOSTEARYL ALCOHOL (UNII: MINERAL OIL (UNII: T5L8T28FGP)	5P) 9Q167V3) II9T) 91C86K) FE (UNII: 7ZBS06BH4B) :1OH) 2DMT128M1S)			S	trength
CETEARETH-12 (UNII: 7V4MR24V PROPYLENE GLYCOL (UNII: 6DC METHYLPARABEN (UNII: A2I8C7F EDETATE DISODIUM (UNII: 7FLD TROLAMINE (UNII: 903K93S3TK) SODIUM CETOSTEARYL SULFA PROPYLPARABEN (UNII: Z8IX2SC WATER (UNII: 059QF0K00R) CETOSTEARYL ALCOHOL (UNII: MINERAL OIL (UNII: T5L8T28FGP) .ALPHATOCOPHEROL (UNII: H4	5P) 9Q167V3) II9T) 91C86K) FE (UNII: 7ZBS06BH4B) 10H) 2DMT128M1S) N855PNZ1)			S	trength
CETEARETH-12 (UNII: 7V4MR24V PROPYLENE GLYCOL (UNII: 6DC METHYLPARABEN (UNII: A218C7F EDETATE DISODIUM (UNII: 7FLD TROLAMINE (UNII: 903K93S3TK) SODIUM CETOSTEARYL SULFA PROPYLPARABEN (UNII: Z81X2SC WATER (UNII: 059QF0K00R) CETOSTEARYL ALCOHOL (UNII: MINERAL OIL (UNII: T5L8T28FGP)	5P) 9Q167V3) II9T) 91C86K) FE (UNII: 7ZBS06BH4B) 10H) 2DMT128M1S) N855PNZ1) P)			S	trength

μ	roduct Chara	icteristics					
Color		white	Score				
Shape				Size			
Flavor				Imprint Code			
Contains							
Packaging							
#	ltem Code	Package Description		Marketing Start Date	Marketing End Date		
1	NDC:80489-088- 01	1 in 1 CARTON		01/01/2021			
1		28.3 g in 1 TUBE; Type 0: Not a Combination Product					
2	NDC:80489-088- 02	1 in 1 CARTON			01/01/2021		
		49.5 g in 1 TUB	E; Type 0: Not a Co	ombination			

Marketing	Information
marketing	mation

Marketing	Application Number or Monograph	Marketing Start	Marketing End
Category	Citation	Date	Date
OTC Monograph Drug	M016	01/01/2021	

Labeler - OL PHARMA TECH,LLC (021170377)

Registrant - OL PHARMA TECH, LLC (021170377)

Establishment

Name	Address	ID/FEI	Business Operations
OL PHARMA TECH,LLC (Drs. pharmacy)		021170377	manufacture(80489-088)

Revised: 1/2024

OL PHARMA TECH,LLC