HYDROCORTISONE ACETATE- hydrocortisone acetate suppository Patrin Pharma Inc.

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

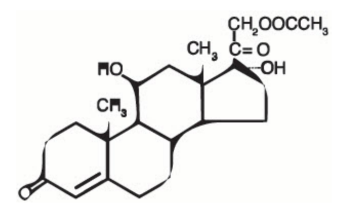
Hydrocortisone Acetate Suppositories

Rx Only

DESCRIPTION

Each Hydrocortisone Acetate Suppository for rectal administration contains hydrocortisone acetate USP in a hydrogenated palm kernel oil base.

Hydrocortisone acetate is a corticosteroid. T he molecular weight of hydrocortisone acetate is 404.50. Chemically, hydrocortisone acetate is pregn-4-ene-3, 20 dione, 21-(acetyloxy)-11, 17-dihydroxy-, (11 β)- with an empirical formula of C₂₃ H₃₂O₆ and the following structural formula:



CLINICAL PHARMACOLOGY

In normal subjects, about 26 percent of hydrocortisone acetate is absorbed when the hydrocortisone acetate suppository is applied to the rectum. Absorption of hydrocortisone acetate may vary across abraded or inflamed surfaces.

Topical steroids are primarily effective because of their anti-inflammatory, antipruritic and vasoconstrictive action.

INDICATIONS AND USAGE

Hydrocortisone Acetate Suppositories are indicated for use in inflamed hemorrhoids, post irradiation (factitial) proctitis, as an adjunct in the treatment of chronic ulcerative colitis, cryptitis, other inflammatory conditions of the anorectum, and pruritus ani.

CONTRAINDICATIONS

Hydrocortisone Acetate Suppositories are contraindicated in those patients with a history of hypersensitivity to any of the components.

PRECAUTIONS

Do not use unless adequate proctologic examination is made.

If irritation develops, the product should be discontinued, and appropriate therapy instituted.

In the presence of an infection, the use of an appropriate antifungal or antibacterial agent should be instituted. If a favorable response does not occur promptly, Hydrocortisone Acetate Suppositories should be discontinued until the infection has been adequately controlled.

Carcinogenesis

No long-term studies in animals have been performed to evaluate the carcinogenic potential of corticosteroid suppositories.

PREGNANCY CATEGORY C

In laboratory animals, topical steroids have been associated with an increase in the incidence of fetal abnormalities when gestating females have been exposed to rather low dosage levels. There are no adequate and well-controlled studies in pregnant women.

Hydrocortisone acetate suppositories should only be used during pregnancy if the potential benefit justifies the risk to the fetus. Drugs of this class should not be used extensively on pregnant patients, in large amounts, or for prolonged periods of time.

It is not known whether this drug is excreted in human milk. Because many drugs are excreted in human milk and because of the potential for serious adverse reactions in nursing infants from hydrocortisone acetate suppositories, a decision should be made whether to discontinue nursing or to discontinue the drug, taking into account the importance of the drug to the mother.

ADVERSE REACTIONS

The following local adverse reactions have been reported with Hydrocortisone Acetate Suppositories: burning, itching, irritation, dryness, folliculitis, hypopigmentation, allergic contact dermatitis, and secondary infection.

To report SUSPECTED ADVERSE REACTIONS, contact Patrin Pharma at 1-800-936-3088 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

DRUG ABUSE AND DEPENDENCE

Drug abuse and dependence have not been reported in patients treated with Hydrocortisone Acetate Suppositories.

OVERDOSAGE

If signs and symptoms of systemic overdosage occur, discontinue use.

DOSAGE AND ADMINISTRATION

For rectal administration: Insert one suppository in the rectum twice daily, morning and night for two weeks, in nonspecific proctitis.In more severe cases, one suppository three times a day or two suppositories twice daily. In factitial proctitis, the recommended duration of therapy is six to eight weeks or less, according to the response of the individual case.

Detach and hold one suppository upright(point upward). Separate tabs at top opening and pull downward to almost the full length of the suppository. Carefully remove the suppository, avoiding excessive handling, which is designed to melt at body temperature. Insert suppository into the rectum, pointed end first, with gentle pressure.

HOW SUPPLIED

25mg (12 count) NDC 39328-029-12 25mg (24 count) NDC 39328-029-24 30mg (12 count) NDC 39328-129-12

Store at 20° to 25°C (68° to 77°F) [see USP Controlled Room Temperature]. Store away from heat. Protect from freezing.

Manufactured For: Patrin Pharma, Skokie, IL 60076 Questions? Call (800) 936 3088

Rev 01.0621

PRINCIPAL DISPLAY PANEL - 25 mg Suppository Carton

NDC 39328-029-12

Rx Only

Hydrocortisone Acetate Suppositories

25 mg

FOR RECTAL USE ONLY

12 Suppositories

PATRIN PHARMA



PRINCIPAL DISPLAY PANEL - 30 mg Suppository Carton

NDC 39328-129-12

Rx Only

Hydrocortisone Acetate Suppositories

30 mg

FOR RECTAL USE ONLY

12 Suppositories

PATRIN PHARMA



HYDROCORTISONE ACETATE

hydrocortisone acetate suppository

Product Information												
Product Type			HUMAN PRESCRIPTION DRUG	Item Code (Source)		NDC:39328-029						
Route of Administration		stration	RECTAL									
_												
A	Active Ingredient/Active Moiety											
		-	edient Name		Basis of Strength		Strength					
HYDROCORTISONE ACETATE (U UNII:W4X0X7BPJ)			VII: 3X7931PO74) (HYDROCORTISONE -		HYDROCORTISONE ACETATE		25 mg					
Ir	nactive Ingre	dients										
			Ingredient Name			Strength						
H	DROGENATED P	PALM OIL (UNII	: 257THB963H)									
D	roduct Chara	octoristics										
	olor		ite to Off-White)		Score							
		BULLET			Size							
	avor				mprint Code							
Contains												
Packaging												
#	ltem Code	Pa	ckage Description		eting Start Date		ting End ate					
1	NDC:39328-029- 12	12 in 1 CARTC Product	N; Type 0: Not a Combination	01/01/202	01/01/2022							
2	NDC:39328-029-	24 in 1 CARTC	N; Type 0: Not a Combination	01/01/202	01/01/2022							
	24	Product										
N	larketing	Informat	ion									
	-				ukating Ctaut	Maula	ting Fud					
Marketing Appl Category		Арриса	ation Number or Monograph Mark Citation		ceting Start Marketin Date Dat		Date					
UNAPPROVED DRUG			01/01/2		2022							
0	ΓHER											
HYDROCORTISONE ACETATE												
hydrocortisone acetate suppository												
пу	diocontisone d	cetate supp	USILOTY									
Product Information												
P	roduct Type		HUMAN PRESCRIPTION DRUG	Item C	ode (Source)	NDC:3	39328-129					

Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:39328-12
Route of Administration	RECTAL		

Active Ingredient/Active Moiety								
		Ingredient Name	Ingredient Name		trength	Strength		
	/DROCORTISONI NII:W4X0X7BPJ)	ACETATE (UNII: 3X7931PO74) (HYDROCORTISC	CETATE (UNII: 3X7931PO74) (HYDROCORTISONE -		ONE	30 mg		
Inactive Ingredients								
Ingredient Name					Strength			
H١	DROGENATED P	ALM OIL (UNII: 257THB963H)						
Product Characteristics								
Co	olor	WHITE (White to Off-White)	S	Score				
Shape		BULLET	Si	Size				
Flavor			In					
Contains								
Packaging								
#	ltem Code	Package Description	Marketing Start Date		Marketing End Date			
1	NDC:39328-129- 12	12 in 1 CARTON; Type 0: Not a Combination Product	01/01/2022					
Marketing Information								
	Marketing Category	Application Number or Monograp Citation	h Mark	eting Start Date		ing End Ite		
UNAPPROVED DRUG OTHER			01/01/2	01/01/2022				

Labeler - Patrin Pharma Inc. (806841677)

Revised: 12/2021

Patrin Pharma Inc.