HARTZ GROOMERS BEST- hydrocortisone spray The Hartz Mountain Corporation

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.

Drug Facts

Active Ingredient

0.5% Hydrocortisone, USP

Purpose

Anti-itch

Uses to soothe • Itching •Flea bites •Dermatitis •Skin Irratations •Eczema

Warnings

For external use on adult dogs and cats only.

Keep out of the reach of children and animnals to avoid unintended consumption.

Do not use •In eyes or nose •not for prolonged use •do not apply to large areas of the body •do not use where infection (pus) is present, since the drug may allow the infection to be spread •on pregnant animals • if product appears to be tasmpered with

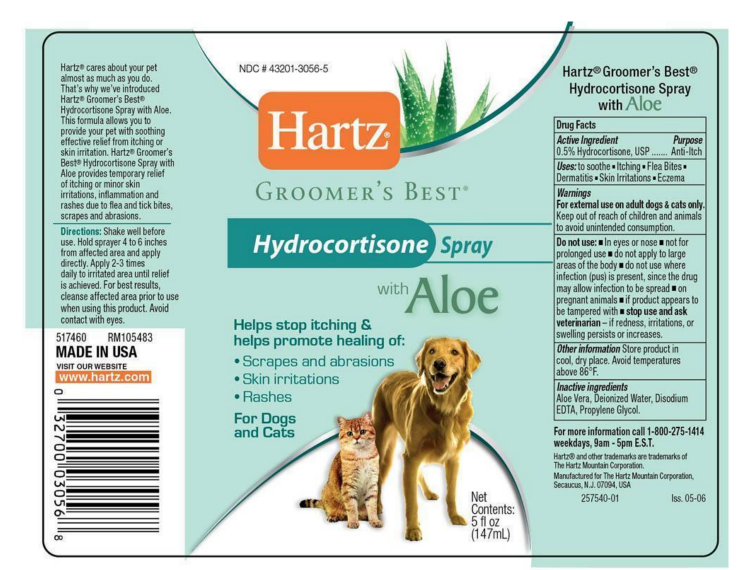
stop use and ask a veterinarian if redness, irritations, or swelling persists or increases

Directions: Shake well before use. Hold sprayer 4 to 6 inches from affected area and apply directly. Apply 2-3 times daily to irritated area until relief is achieved. For best results, cleanse affected area prior to use when using this product. Avoid contact with eyes

*Other information*Store product in cool, dry place. Avoid temperatures above 86F

Inactive Ingredients

Aloe Vera Gel, Deionized Water, Disodium EDTA, Propylene Glycol



HARTZ GROOMERS BEST

hydrocortisone spray

Product Information						
Product Type	OTC ANIMAL DRUG LABEL	Item Code (Source)	NDC:43201-3056			
Route of Administration	TOPICAL	DEA Schedule				

Active Ingredient/Active Moiety						
Ingredient Name	Basis of Strength	Strength				
HYDRO CORTISONE (HYDROCORTISONE)	HYDROCORTISONE	0.5 g in 100 mL				

Inactive Ingredients					
Ingredient Name	Strength				
ALOE VERA LEAF					
WATER					
EDETATE DISODIUM					
PROPYLENE GLYCOL					

Packaging								
#	Item Code	Package Description	Market	ing Start Date	Marketing End Date			
1	NDC:43201-3056-5	147 mL in 1 BOTTLE, SPRAY						
Marketing Information								
N	Marketing Category	Application Number or Monograp	h Citation	Marketing Start Da	te Marketing End Date			

$\pmb{Labeler} \textbf{ - } \textbf{ The Hartz Mountain Corporation (058109158)}$

unapproved drug other

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