

HEMOPROPIN HEMORRHOIDAL- lanolin, petrolatum ointment

Bee Right LLC

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.

Hemopropin Hemorrhoidal Ointment

Drug Facts

Active Ingredient

Lanolin 17.76%

Petrolatum 71.2%

Purpose

Hemorrhoidal Protectant

Uses

- helps relieve itching and discomfort associated with hemorrhoids and inflamed hemorrhoidal tissues
- temporarily protects irritated areas
- temporarily relieves burning
- temporarily protects inflamed, irritated anorectal surface to help make bowel movements less painful

Warnings

For rectal use only.

When using this product

do not exceed the recommended daily dosage unless directed by doctor.

Stop use and consult a doctor if:

- condition worsens or does not improve within 7 days
- the introduction of the applicator into the rectum causes additional pain or bleeding occurs

If pregnant or breastfeeding,

consult a health care professional before use.

Keep out of reach of children.

If swallowed, get medical help or contact a Poison Control Center right away.

Directions

- Adults: when practical, cleanse the affected area with mild soap and warm water and rinse thoroughly; gently dry by patting or blotting with toilet tissue or a soft cloth before application of this product
- Apply externally to the affected area up to 6 times daily or after each bowel movement
- FOR INTRARECTAL USE: attach applicator to tube; lubricate applicator well, then gently insert

applicator into the rectum

- Children under 12 years of age: consult a doctor

Other information

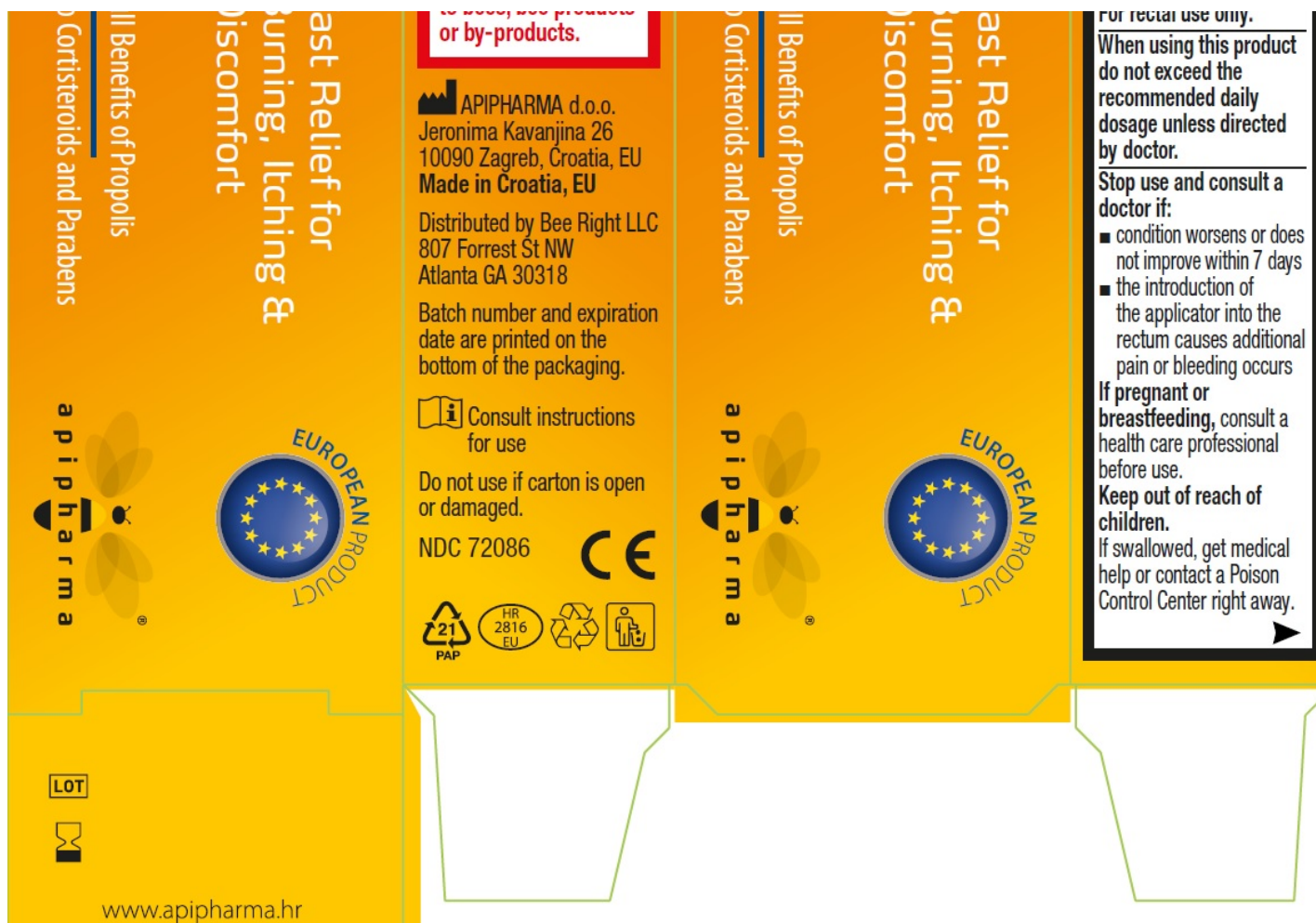
- store in a dry and cool place

Inactive ingredients

Alcohol, propolis wax, roman chamomile, water

Package Labeling:






Drug Facts **Active Ingredients** Lanolin 17.76%, Petrolatum 71.2% **Purpose** Extra Strength Hemorrhoidal Protectant **Uses** ■ helps relieve itching and discomfort associated with hemorrhoids and inflamed hemorrhoidal tissues ■ temporarily protects irritated areas ■ temporarily relieves burning ■ temporarily protects inflamed, irritated anorectal surface to help make bowel movements less painful **Warnings** **For rectal use only.** When using this product do not exceed the recommended daily dosage unless directed by doctor. **Stop use and consult a doctor if:** ■ condition worsens or does not improve within 7 days ■ the introduction of the applicator into the rectum causes additional pain or bleeding occurs **If pregnant or breastfeeding,** consult a health care professional before use. **Keep out of reach of children.** If swallowed, get medical help or contact a Poison Control Center right away.

REF 1101 US-10     2460  

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Hemopropin[®]

Extra Strength
Hemorrhoidal Ointment

 Net Wt. 0.7 oz (20g)

Fast Relief
for Burning,
Itching &
Discomfort

HEMOPROPIN HEMORRHOIDAL

lanolin, petrolatum ointment

Product Information

Product Type

HUMAN OTC DRUG

Item Code (Source)

NDC:71952-010

| | | | | |
|--|--|---|-----------------------------|---------------------------|
| Route of Administration | TOPICAL | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | LANOLIN (UNII: 7EV65EAW6H) (LANOLIN - UNII:7EV65EAW6H) | LANOLIN | 17.76 g in 100 g | |
| | PETROLATUM (UNII: 4T6H12BN9U) (PETROLATUM - UNII:4T6H12BN9U) | PETROLATUM | 71.2 g in 100 g | |
| Inactive Ingredients | | | | |
| | Ingredient Name | Strength | | |
| | ALCOHOL (UNII: 3K9958V90M) | | | |
| | PROPOLIS WAX (UNII: 6Y8XYV2NOF) | | | |
| | CHAMAEMELUM NOBILE (UNII: 7NF3GE7IWR) | | | |
| | WATER (UNII: 059QF0KO0R) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:71952-010-00 | 1 in 1 BOX | 07/01/2018 | |
| 1 | | 20 g in 1 TUBE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| OTC monograph final | part346 | 07/01/2018 | | |

Labeler - Bee Right LLC (080988303)

Revised: 7/2018

Bee Right LLC