PRORINO DELAY- menthol, eucalyptus globulus, mentha arvensis cream HOT PRODUCTIONS AND VERTRIEBS GMBH

Disclaimer: This homeopathic product has not been evaluated by the Food and Drug Administration for safety or efficacy. FDA is not aware of scientific evidence to support homeopathy as effective.

HOT PRODUCTS - Prorino Delay Cream (71326-102)

ACTIVE INGREDIENTS

Menthol 1x HPUS

Eucalyptus Globulus Leaf Oil 0.1X HPUS

Mentha Arvenses Leaf Oil 0.05x HPUS

PURPOSE

Topical Anesthetic

USE

For temporary male genital desensitization, helping to slow the onset of ejaculation.

WARNINGS

Premature ejaculation may be due to a condition requiring medical supervision. If this product, used as directed, does not provide relief, discontinue use and consult a doctor. Avoid contact with the eyes or broken skin.

If you or your partner develop a rash or irritation, such as burning or itching, discontinue use. If symptoms persist, consult a doctor.

Do not use during the partner's pregnancy.

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

DIRECTIONS

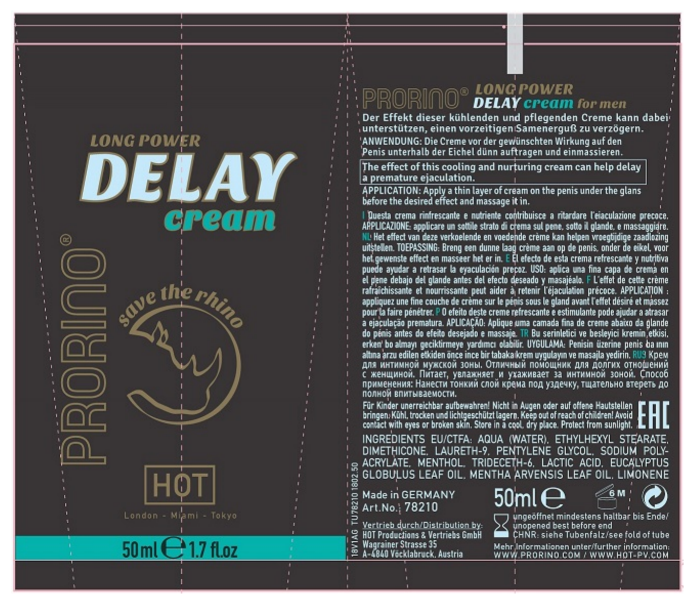
- Prior to the coitus apply a thin layer of the cream onto the glans and under it (glans frenulum) and spread by massaging.
- Approximately 5 minutes later the cream will have absorbed well and will have fully developed its effect.
- Prior to the coitus itself, the cream must be unconditionally washed off the penis and glans.
- The effect may be increased by repeated use of the cream shortly before the coitus.

OTHER INFORMATION

• Store in a cool, dry place. Protect from sunlight.

INACTIVE INGREDIENTS

Water (Aqua), Ethylhexyl Stearate, Dimethicone, Laureth-9, Pentylene Glycol, Sodium Polyacrylate, Trideceth-6, Lactic Acid, Limonene



Drug Facts

Active Ingredients

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PRORINO DELAY

menthol, eucalyptus globulus, mentha arvensis cream

Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:71326-102 Route of Administration TOPICAL

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
MENTHOL (UNII: L7T10EIP3A) (MENTHOL - UNII:L7T10EIP3A)	MENTHOL	1 g in 100 mL		
EUCALYPTUS OIL (UNII: 2R040NI662) (EUCALYPTUS OIL - UNII:2R040NI662)	EUCALYPTUS OIL	0.097 g in 100 mL		
MENTHA ARVENSIS LEAF OIL (UNII: 1AEY1M553N) (MENTHA ARVENSIS LEAF OIL - UNII:1AEY1M553N)	MENTHA ARVENSIS LEAF OIL	0.0478 g in 100 mL		

Inactive Ingredients				
Ingredient Name	Strength			
WATER (UNII: 059QF0KO0R)				
ETHYLHEXYL STEARATE (UNII: EG3PA2K3K5)				
DIMETHICONE (UNII: 92RU3N3Y1O)				
POLIDOCANOL (UNII: 0AWH8BFG9A)				
PENTYLENE GLYCOL (UNII: 50C1307PZG)				
SODIUM POLYACRYLATE (8000 MW) (UNII: 285CYO341L)				
TRIDECETH-6 (UNII: 3T5PCR2H0C)				
LACTIC ACID (UNII: 33X04XA5AT)				
LIMONENE, (+/-)- (UNII: 9MC3I34447)				

P	Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:71326-102- 51	1 in 1 BOX	08/29/2018			
1	NDC:71326-102- 11	50 mL in 1 TUBE; Type 0: Not a Combination Product				

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
unapproved homeopathic		08/29/2018		

Labeler - HOT PRODUCTIONS AND VERTRIEBS GMBH (300011984)

Revised: 10/2023 HOT PRODUCTIONS AND VERTRIEBS GMBH