TIGER DELAYSPRAY- lidocaine spray Magverz INC

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.

TIGER Spray

Lidocaine HCI 10 % ... External Anesthetic

Purpose

External Anesthetic

Keep out of Reach of Children

WARNINGS: Keep out of reach of children. For External use only

Indication and Usage

temporarily reduces sensitivity of the penis, which helps to delay ejaculation in cases of over-rapid or premature ejaculation (coming to a climax too quickly)

Inactive Ingredients

Warning

Allergy alert: do not use this product if you or your partner are allergic (sensitive) to local Anesthetics.

Do not use - on broken or inflamed skin - if your partner is pregnant

Ask a doctor before use if you have, or ever had, liver or kidney problems

Ask a doctor or pharmacist before use if you are already taking prescribed drugs

when using this product

- do not get into eyes
- do not inhale
- do not exceed 24 sprays in 24 hours

Dosage and Administration

-apply 3 or more sprays, not to exceed 10, to the head and shaft of the penis before intercourse, or as directed by a doctor

-wash product after intercourse

-correct quantity and time of application will be determined by individual requirements and you should always use the minimum effective quantity

Product Label



TIGER DELAYSPRAY									
lidocaine spray									
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Product Information									
Product T ype	HUMAN OTC DRUG	Item Code (Source)		NDC:59240-005					
Route of Administration	TOPICAL								
Active Ingredient/Active Moiety									
Ingredient Name			Basis of Strengt	h	Strength				
LIDO CAINE (UNII: 98PI200987) (LIDO CAINE - UNII:98PI200987)			LIDOCAINE		10 mg in 100 mL				

I	nactive Ingredi	ents		
		Strength		
IS	SOPROPYL MYRIS	FATE (UNII: 0 RE8 K4LNJS)		
S	TEARIC ACID (UNII	: 4ELV7Z65AP)		
.A	ALPHATO COPHE	ROL (UNII: H4N855PNZ1)		
P	ackaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:59240-005- 06	6 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	0 1/0 3/20 17	
2	NDC:59240-005- 08	8 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	0 1/0 3/20 17	
3	NDC:59240-005- 12	12 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	0 1/0 3/20 17	
	Aarketing In	formation		
N		Ann liestion Number or Meneguenh Citation	Marketing Start Date	Marketing End Date
	Marketing Catego	ry Application Number or Monograph Citation	inter ine ting o tart Date	

Labeler - Magverz INC (078712269)

Registrant - Magverz INC (078712269)

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Magverz INC