# ECLIPSE TOPICAL ANALGESIC LL- tetracaine hcl cream Sambria Pharmaceuticals, 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.

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## **Eclipse LL Topical Analgesic**

## **Active Ingredients**

Tetracaine HCL 2.0% w/w

## **□**Purpose

External Analgesic

#### $\Box$ Uses

For temporary relief of pain and itching due to minor skin irritation

## **IFor external use only**

## **Avoid contact with eyes**

**□Do not use** in large quantities, particularly over raw surfaces or blistered areas

## **Stop use and ask a doctor if**■

• Condition worsens, or if symptoms persist for more then 7 days or clear up and occur again within a few days. Discontinue use.

#### **Example 2 IXeep out of reach of children**

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

#### Directions

For adults and children two-years or older: Apply to affected area not more than 3 to 4 times daily. Children under 2 years of age: consult a physician.

## **Inactive Ingredients**

Aqua (Deionized Water), Arnica Montana Flower Extract, C13-14 Isoparaffin, Chondroitin Sulfate, Emu Oil, Ethoxydiglycol, Ethylhexylglycerin, Glucosamine Sulfate, Isopropyl Palmitate, Laureth-7, Melaleuca Alternifolia (Tea Tree) Leaf Oil, Methylsulfonylmethane (MSM), Phenoxyethanol, Polyacrylamide, Propylene Glycol, Stearic Acid, Triethanolamine

#### **Other Information**

Protect this product from excessive heat and direct sun.

## **Questions or Comments?**

FDA Registered: NDC No. 54723-667-03

800-759-6876



## ECLIPSE TOPICAL ANALGESIC LL

tetracaine hcl cream

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:54723-669
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
TETRACAINE HYDRO CHLO RIDE (UNII: 5NF5D4OPCI) (TETRACAINE - UNII: 06 19 F35CGV)	TETRACAINE HYDROCHLORIDE	20 mg in 1000 mg	

Inactive Ingredients			
Ingredient Name	Strength		
WATER (UNII: 059QF0KO0R)			
ARNICA MONTANA FLOWER (UNII: OZ0E5Y15PZ)			
C13-14 ISOPARAFFIN (UNII: E4F12ROE70)			
CHONDROITIN SULFATE SODIUM (BOVINE) (UNII: 8QTV3DTT8W)			
EMU O IL (UNII: 344821WD61)			
DIETHYLENE GLYCOL MONOETHYL ETHER (UNII: A1A118 X02B)			
ETHYLHEXYLGLYCERIN (UNII: 147D247K3P)			
GLUCO SAMINE SULFATE (UNII: 1FW7WLR731)			
ISOPROPYL PALMITATE (UNII: 8CRQ2TH63M)			
<b>LAURETH-7</b> (UNII: Z95S6G8201)			

MELALEUCA ALTERNIFO LIA LEAF (UNII: G43C57162K)		
DIMETHYL SULFONE (UNII: 9H4PO4Z4FT)		
PHENO XYETHANOL (UNII: HIE492ZZ3T)		
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)		
STEARIC ACID (UNII: 4ELV7Z65AP)		
TROLAMINE (UNII: 9O3K93S3TK)		

Packaging				
#	Item Code	Package Description	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
1	NDC:54723-669-03	3000 mg in 1 PACKET; Type 0: Not a Combination Product	02/01/2016	
Marketing Information				
Marketing Information				

Marketing Start Date

 $0\,2/0\,1/20\,16$ 

Marketing End Date

Application Number or Monograph Citation

## Labeler - Sambria Pharmaceuticals, Inc. (078676259)

Marketing Category

OTC monograph not final part348

Establishment				
Name	Address	ID/FEI	Business Operations	
A.I.G. Technologies, Inc.		171837367	manufacture(54723-669)	

Establishment				
Name	Address	ID/FEI	Business Operations	
JP Packaging LLC		151369456	repack(54723-669)	

Revised: 8/2018 Sambria Pharmaceuticals, Inc.