# NOREPINEPHRINE- norepinephrine (bitartrate) liquid Deseret Biologicals, Inc.

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.

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#### Drug Facts:

### **ACTIVE INGREDIENT:**

Norepinephrine (Bitartrate) 8X, 12X, 30X, 200X, 12C, 30C, 60C, 200C.

#### **HOMEOPATHIC INDICATIONS:**

For temporary relief of symptoms related to Norepinephrine sensitivity including rashes, hives, Premenstrual Syndrome, and headache.\*\*

\*\*These statements are based upon traditional homeopathic principles. They have not been reviewed by the Food and Drug Administration.

#### WARNINGS:

**Keep out of reach of children.** In case of overdose, contact physician or a Poison Control Center right away.

If pregnant or breast-feeding, ask a health professional before use.

Tamper seal: "Sealed for Your Protection." Do not use if seal is broken or missing.

#### **KEEP OUT OF REACH OF CHILDREN:**

**Keep out of reach of children.** In case of overdose, contact physician or a Poison Control Center right away.

#### DIRECTIONS:

1-10 drops under the tongue, 3 times a day or as directed by a health professional. Consult a physician for use in children under 12 years of age.

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#### **INACTIVE INGREDIENTS:**

Demineralized Water, 25% Ethanol

#### **QUESTIONS:**

Dist. By: Deseret Biologicals, Inc. 469 W. Parkland Drive Sandy, UT 84070 www.desbio.com

#### PACKAGE LABEL DISPLAY:

DESBIO NDC 43742-1489-1 HOMEOPATHIC NOREPINEPHRINE 1 FL OZ (30 ml)

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LOT:

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HOMEOPATHIC

NOREPINEPHRINE



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## NOREPINEPHRINE

norepinephrine (bitartrate) liquid

Product Information								
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:43742-1489				
Route of Administration	ORAL							
Active Ingredient/Active Moiety								
Ingredient Name			Basis of Strength		Strength			
<b>NOREPINEPHRINE BITARTRATE</b> (UNII: IFY5PE3ZRW) (NOREPINEPHRINE - UNII:X4W3ENH1CV)					8 [hp_X] in 1 mL			

Inactive Ingredients								
		Strength						
WATER (UNII: 059QF0K00R)								
AL	ALCOHOL (UNII: 3K9958V90M)							
Packaging								
#	ltem Code	Package Description		Marketing Start Ma Date				
1	NDC:43742- 1489-1	30 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product	05/28/2019					
Marketing Information								
	Marketing Category	Application Number or Monograph Citation	Marketing Date	-	Marketing End Date			
	approved		05/28/2019					

## Labeler - Deseret Biologicals, Inc. (940741853)

# Registrant - Apotheca Company (844330915)

Establishment							
Name	Address	ID/FEI	Business Operations				
Apotheca Company		844330915	manufacture(43742-1489) , api manufacture(43742-1489) , label(43742-1489) , pack(43742-1489)				

Revised: 1/2023

Deseret Biologicals, Inc.

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