

PHLORIDZIN PHENOLIC- phloridzinum liquid
Energique, 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.

DRUG FACTS:

ACTIVE INGREDIENTS:

(in each drop): 33.30% of Phloridzinum 30X, 12C, 30C; 0.10% of Phloridzinum 12X.

INDICATIONS:

May temporarily relieve symptoms associated with reactions to Phloridzin such as abnormalities related to sugar.**

**Claims based on traditional homeopathic practice, not accepted medical evidence. Not FDA evaluated.

WARNINGS:

If pregnant or breastfeeding, ask a health professional before use.

Keep out of reach of children. In case of overdose, get medical help or contact a Poison Control Center right away.

Do not use if tamper evident seal is broken or missing.

Store in a cool, dry place.

KEEP OUT OF REACH OF CHILDREN:

Keep out of reach of children. In case of overdose, get medical help or contact a Poison Control Center right away.

DIRECTIONS:

Adults and children 5 to 10 drops orally, 1 time daily or as otherwise directed by a health care professional. If symptoms persist for more than 7 days, consult your health care professional. Consult a physician for use in children under 12 years of age.

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FDA evaluated.

INACTIVE INGREDIENTS:

Demineralized water, 20% Ethanol

QUESTIONS:

Dist. by Energique, Inc.

201 Apple Blvd.

Woodbine, IA 51579 **800.868.8078**

PACKAGE LABEL DISPLAY:

ENERGIQUE

SINCE 1987

HOMEOPATHIC REMEDY

PHLORIDZIN

PHENOLIC

1 fl. oz. (30 ml)

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LOT: XXXXXX

PHLORIDZIN PHENOLIC

phloridzinum liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:44911-0509
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PHLORIZIN (UNII: CU9S17279X) (PHLORIZIN - UNII:CU9S17279X)	PHLORIZIN	12 [hp_X] in 1 mL

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
ALCOHOL (UNII: 3K9958V90M)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:44911-0509-1	30 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product	08/06/2019	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved homeopathic		08/06/2019	

Labeler - Energique, Inc. (789886132)**Registrant** - Apotheca Company (844330915)**Establishment**

Name	Address	ID/FEI	Business Operations
Apotheca Company		844330915	manufacture(44911-0509) , api manufacture(44911-0509) , label(44911-0509) , pack(44911-0509)

Revised: 10/2024

Energique, Inc.