

TOXOPLASMOSIS NOSODE 4034- toxoplasmosis nosode liquid

Professional Complementary Health Formulas

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.

N34

ACTIVE INGREDIENTS

Toxoplasmosis 12X, 30X

QUESTIONS

Professional Formulas

PO Box 2034 Lake Oswego, OR 97035

INDICATIONS

For the temporary relief of muscle pain, occasional headache, or fatigue.*

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

WARNINGS

Severe or persistent symptoms may be a sign of a serious condition. Consult a doctor promptly if symptoms persist or are accompanied by a fever. Keep out of the reach of children. In case of overdose, get medical help or contact a poison control center right away. If pregnant or breastfeeding, ask a healthcare professional before use.

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DIRECTIONS

Place drops under tongue 30 minutes before/after meals. Adults and children 12 years and over: Take 10 to 15 drops once weekly or monthly. If mild symptoms are present, take 10 drops up to 3 times per day. Consult a physician for use in children under 12 years of age.

OTHER INFORMATION

Tamper resistant. If seal is broken, do not use. After opening, close container tightly and store at room temperature away from heat.

INACTIVE INGREDIENTS

20% ethanol, purified water.

LABEL

Est 1985
Professional Formulas
Complementary Health
Toxoplasmosis Nosode
Homeopathic Remedy
2 FL. OZ. (59 mL)



TOXOPLASMOSIS NOSODE 4034

toxoplasmosis nosode liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:63083-4034
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
TOXOPLASMA GONDII (UNII: BMV90JF469) (TOXOPLASMA GONDII - UNII:BMV90JF469)	TOXOPLASMA GONDII	12 [hp_X] in 59 mL

Inactive Ingredients				
Ingredient Name			Strength	
ALCOHOL (UNII: 3K9958V90M)				
WATER (UNII: 059QF0KO0R)				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:63083-4034-2	59 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product	08/15/1985	
Marketing Information				
Marketing Category		Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved homeopathic			08/15/1984	

Labeler - Professional Complementary Health Formulas (167339027)

Registrant - Natural Pharmaceutical Manufacturing LLC (015624923)

Establishment			
Name	Address	ID/FEI	Business Operations
Natural Pharmaceutical Manufacturing LLC		015624923	manufacture(63083-4034)

Revised: 1/2026

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