

**N-13- ceanothus americanus leaf, urtica urens, sodium chloride, lachesis muta whole, cinchona officinalis bark, lycopodium clavatum whole, ferric chloride hexahydrate, and sus scrofa spleen solution
DNA Labs, 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.

N-13

NDC 58264-0211-1

INDICATIONS

Support for the Spleen; Blood and Immune system.

INGREDIENTS

ACTIVE

Ceanothus 4x, Urtica urens 4x, Natrum muriaticum 6x, Lachesis mutus 12x, Cinchona officinalis 8x, Lycopodium clavatum 6x, Ferrum muriaticum 6x, Spleen 6/12/30/60/200x

INACTIVE

20% alcohol in purified water.

SUGGESTED DOSAGE

One dropper under tongue two times daily. Acute symptoms ½ dropper under tongue every 30 minutes for two hours.

SHAKE WELL

Warnings

- Use only if cap seal is unbroken.
- If pregnant or breastfeeding, ask a healthcare professional before use.
- Keep this and all medication out of the reach of children.

To be used according to standard homeopathic indications.

PRINCIPAL DISPLAY PANEL - 1 FL. OZ. Bottle Label

DYNAMIC

NUTRITIONAL

ASSOCIATES, INC.

N-13 SPLEEN ENERGIZER

HOMEOPATHIC ENDOCRINE

1 FL. OZ.

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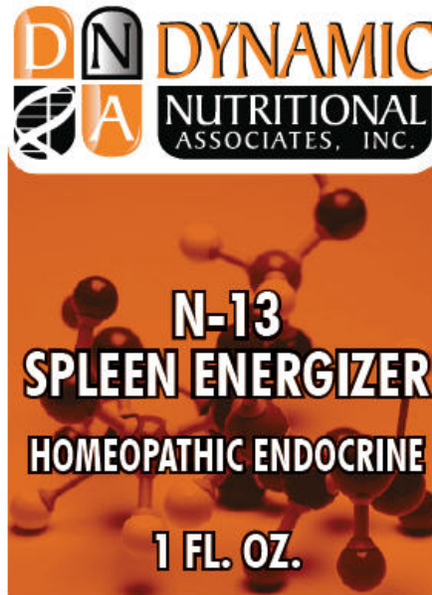
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Rev. 3/18



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MANUFACTURED FOR:

DNA LABORATORIES, INC.
Coeur d'Alene, ID 83814

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Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:58264-0211
Route of Administration	SUBLINGUAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CEANOTHUS AMERICANUS LEAF (UNII: 25B1Y14T8N) (CEANOTHUS AMERICANUS LEAF - UNII:25B1Y14T8N)	CEANOTHUS AMERICANUS LEAF	4 [hp_X] in 1 mL
URTICA URENS (UNII: IHN2NQ5OF9) (URTICA URENS - UNII:IHN2NQ5OF9)	URTICA URENS	4 [hp_X] in 1 mL
SODIUM CHLORIDE (UNII: 451W47IQ8X) (CHLORIDE ION - UNII:Q32ZN48698, SODIUM CATION - UNII:LYR4M0NH37)	SODIUM CHLORIDE	6 [hp_X] in 1 mL
LACHESIS MUTA WHOLE (UNII: 6Y582I290C) (LACHESIS MUTA WHOLE - UNII:6Y582I290C)	LACHESIS MUTA WHOLE	12 [hp_X] in 1 mL
CINCHONA OFFICINALIS BARK (UNII: S003A158SB) (CINCHONA OFFICINALIS BARK - UNII:S003A158SB)	CINCHONA OFFICINALIS BARK	8 [hp_X] in 1 mL
LYCOPODIUM CLAVATUM WHOLE (UNII: 005ICF6L27) (LYCOPODIUM CLAVATUM WHOLE - UNII:005ICF6L27)	LYCOPODIUM CLAVATUM WHOLE	6 [hp_X] in 1 mL

FERRIC CHLORIDE HEXAHYDRATE (UNII: 0I2XIN602U) (FERRIC CATION - UNII:91O4LML611)	FERRIC CATION	6 [hp_X] in 1 mL
SUS SCROFA SPLEEN (UNII: 92AMN5J79Y) (SUS SCROFA SPLEEN - UNII:92AMN5J79Y)	SUS SCROFA SPLEEN	200 [hp_X] in 1 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:58264-0211-1	29.57 mL in 1 BOTTLE, GLASS; Type 0: Not a Combination Product	01/01/1990	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved homeopathic		01/01/1990	

Labeler - DNA Labs, Inc. (031784339)

Revised: 1/2025

DNA Labs, Inc.