

**D-46- ferrosferric phosphate, lithium carbonate, sodium sulfate, strychnos nux-vomica seed, rhododendron aureum leaf, and filipendula ulmaria root 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.*

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**D-46**

**NDC 58264-0046-1**

**INDICATIONS**

Rheumatism, gout, swollen joints.

**INGREDIENTS**

**ACTIVE**

Ferrum phosphoricum 12x, Lithium carbonicum 12x, Natrum sulfuricum 30x, Nux vomica 30x, Rhododendron chrysanthum 6x, Spiraea ulmaria 12x

**INACTIVE**

20% alcohol in purified water.

**Rx CAUTION**

Federal law prohibits dispensing without a prescription.

**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.

## D 46

## HOMEOPATHIC

## STRESS FORMULA

## 1 FL. OZ.

### SUGGESTED DOSAGE:

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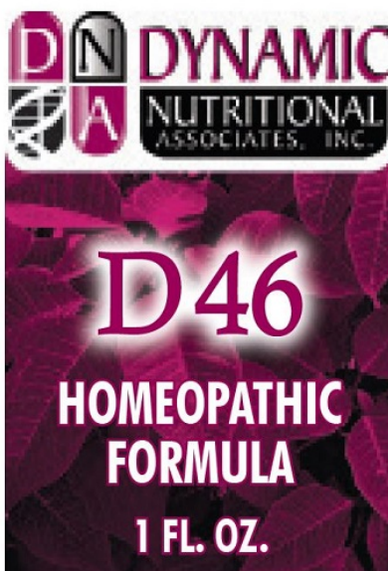
### SHAKE WELL

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Rev. 12/22



NDC 58264-0046-1

### INDICATIONS:

Rheumatism, gout, swollen joints.

### INGREDIENTS:

**ACTIVE:** Natrum sulfuricum, Nux vomica 30x, Ferrum phosphoricum, Lithium carbonicum, Spiraea ulmaria 12x, Rhododendron chrysanthum 6x

**INACTIVE:** 20% alcohol in purified water.

### Rx CAUTION:

Federal law prohibits dispensing without a prescription.

Mfg for: **DNA LABORATORIES, INC.**  
Chelan, WA 98816

## D-46

ferrosoferric phosphate, lithium carbonate, sodium sulfate, strychnos nux-vomica seed, rhododendron aureum leaf, and filipendula ulmaria root solution

### Product Information

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:58264-0046
<b>Route of Administration</b>	SUBLINGUAL		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>FERROSFERRIC PHOSPHATE</b> (UNII: 91GQH8I5F7) (FERROSFERRIC PHOSPHATE - UNII:91GQH8I5F7)	FERROSFERRIC PHOSPHATE	12 [hp_X] in 1 mL
<b>LITHIUM CARBONATE</b> (UNII: 2BMD2GNA4V) (LITHIUM CATION - UNII:8H8Z5UER66)	LITHIUM CARBONATE	12 [hp_X] in 1 mL
<b>SODIUM SULFATE</b> (UNII: 0YPR65R21J) (SODIUM SULFATE ANHYDROUS - UNII:36KCS0R750)	SODIUM SULFATE	30 [hp_X] in 1 mL
<b>STRYCHNOS NUX-VOMICA SEED</b> (UNII: 269XH13919) (STRYCHNOS NUX-	STRYCHNOS NUX-	30 [hp_X]

VOMICA SEED - UNII:269XH13919)	VOMICA SEED	in 1 mL
<b>RHODODENDRON AUREUM LEAF</b> (UNII: IV92NQJ73U) (RHODODENDRON AUREUM LEAF - UNII:IV92NQJ73U)	RHODODENDRON AUREUM LEAF	6 [hp_X] in 1 mL
<b>FILIPENDULA ULMARIA ROOT</b> (UNII: 997724QNDS) (FILIPENDULA ULMARIA ROOT - UNII:997724QNDS)	FILIPENDULA ULMARIA ROOT	12 [hp_X] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>ALCOHOL</b> (UNII: 3K9958V90M)	
<b>WATER</b> (UNII: 059QF0KO0R)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:58264-0046-1	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.