#### PHOSPHORUS- phosphorus liquid Newton Laboratories, 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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### Phosphorus 6417L

## **INDICATIONS & USAGE SECTION**

Fever; Infection; Irritability; Sore Throat; Cough; Indigestion; Sluggish elimination

## **DOSAGE & ADMINISTRATION SECTION**

**Directions:** Ages 12 and up, take 6 drops by mouth (ages 0 to 11, give 3 drops) as needed or as directed by a health professional. Sensitive persons begin with 1 drop and gradually increase to full dose.

## **OTC - ACTIVE INGREDIENT SECTION**

Phosphorus 15x, 10x, 200c, 30c.

## **OTC - PURPOSE SECTION**

Fever; Infection; Irritability; Sore Throat; Cough; Indigestion; Sluggish elimination

## **INACTIVE INGREDIENT SECTION**

**Inactive Ingredients**: USP Purified water; USP Gluten-free, non-GMO, organic cane alcohol 20%.

## **QUESTIONS SECTION**

newtonlabs.net - Questions? 1.800.448.7256 Newton Laboratories, Inc. FDA Est # 1051203 - Conyers, GA 30013

## WARNINGS SECTION

**WARNINGS:Keep out of reach of children. Do not use** if tamper-evident seal is broken or missing. If symptoms worsen or persist for more than a few days, consult a doctor. If **pregnant or breast-feeding**, ask a doctor before use.

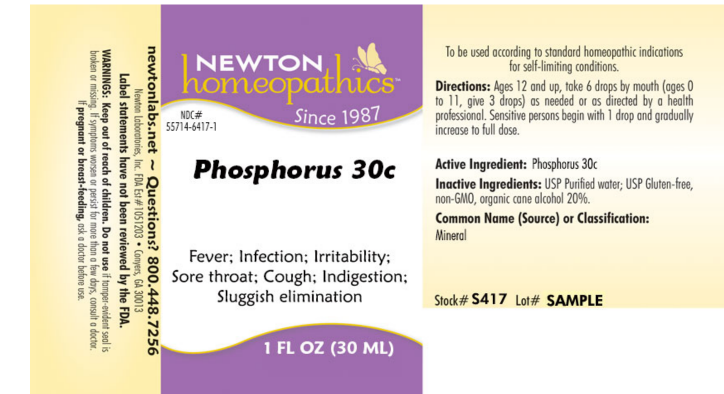
## **OTC - PREGNANCY OR BREAST FEEDING SECTION**

If **pregnant or breast-feeding**, ask a doctor before use.

## **OTC - KEEP OUT OF REACH OF CHILDREN SECTION**

Keep out of reach of children.

PACKAGE LABEL



PHOSPHORUS						
phosphorus liquid						
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<b>Product Information</b>						
Product Type	HUMAN OTC DRUG	Item Code (Source)		ND	NDC:55714-6417	
Route of Administration	ORAL					
Active Ingradiant/Active Main	A#7					
Active Ingredient/Active Moiety Ingredient Name Basis of					Strength	
		PHOSPHORU		15 [hp_X] in 1 mL		
Inactive Ingredients						
Ingredient Name				Strength		
ALCOHOL (UNII: 3K9958V90M)						
WATER (UNII: 059QF0KO0R)						
Packaging						
# Item Code	Package Description		Marketing Date	Start	Marketing End Date	
1 NDC:55714-6417- 30 mL in 1 BOTTL 1 Product	E, GLASS; Type 0: Not a Combi	nation 09	9/01/2011			

Marketing Information						
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
unapproved homeopathic		09/01/2011				
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# Labeler - Newton Laboratories, Inc. (788793610)

Registrant - Newton Laboratories, Inc. (788793610)

# Establishment

Name	Address	ID/FEI	<b>Business Operations</b>
Newton Laboratories, Inc.		788793610	manufacture(55714-6417)

Revised: 12/2020

Newton Laboratories, Inc.