KALI MURIATICUM - kali muriaticum 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.

Kali muriaticum

## INDICATIONS \& USAGE SECTION

Cell Salt; Nose, throat \& ear discharge; Dandruff; Loss of voice; Hemorrhoids, Swelling around joints.

## 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

Kali muriaticum 15x, 10x, 200c, 30c.

## OTC - PURPOSE SECTION

Cell Salt; Nose, throat \& ear discharge; Dandruff; Loss of voice; Hemorrhoids, Swelling around joints.

## INACTIVE INGREDIENT SECTION

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

## QUESTIONS SECTION

www.newtonlabs.net Newton Laboratories, Inc. FDA Est \# 1051203 - Conyers, GA 30012
Questions? 1.800.448.7256

## 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


## NEWTON

## 

Cell Salt; Nose, throat and ear discharge; Dandruff; loss of voice; Hemorthoids; Swelling around joints

1 fl oz (29.57 mil)

## KALI MURIATICUM

kali muriaticum liquid

## Product Information

Product Type
Route of Administration

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HUMAN OTC DRUG Item Code (Source)
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NDC:55714-6297

Active Ingredient/Active Moiety

|  | Ingredient Name | Basis of Strength | Strength |
| :--- | :--- | :--- | :--- |
| Potassium Chloride (UNII: 660 YQ98110) (Chloride Ion - UNII:Q32ZN48698) | Potassium Chloride | $15[\mathrm{hp}$ _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 |
| :--- | ---: | ---: | ---: | ---: |

## Marketing Information

Marketing Category Application Number or Monograph Citation Marketing Start Date Marketing End Date
unapproved homeopathic 09/0 1/2011

Labeler - Newton Laboratories, Inc. (788793610)

Registrant - Newton Laboratories, Inc. (788793610)

## Establishment

Name
Newton Laboratories, Inc.

Address
ID/FEI
788793610

