

RENAKARE- potassium gluconate powder
Neogen Corporation-Mercer Rd

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

RenaKare™

Potassium Gluconate Powder

CAUTION: Federal law restricts this drug to use by or on the order of a licensed veterinarian.

FOR ANIMAL USE ONLY

Each level ¼ teaspoon (0.65 g) contains (minimum):

Potassium gluconate.....2 mEq (468 mg)

in a palatable base

Dosage:

Adult cats and dogs.....2 mEq/4.5 kg (¼ tsp/10 lb) PO BID

Kittens and puppies.....Consult veterinarian

Administer with food. Dosage may be adjusted as necessary by the veterinarian.

Store in a dry place at controlled room temperature 15°-30°C (59°-86°F). Keep container tightly closed.

Item No. 09075

KEEP OUT OF REACH OF CHILDREN

Indications:

For use as a potassium supplement in hypokalemic cats and dogs.

Warnings:

May cause hyperkalemia. Do not exceed the recommended dose. Frequent patient monitoring is advised. Adverse effects include gastrointestinal irritation, muscular weakness, twitching, irritability, and cardiac conduction disturbances. Potassium salts are contraindicated in animals with hyperkalemia, severe renal impairment or failure, severe hemolytic reactions, Addison's disease, decreased gastrointestinal motility, and acute dehydration.

Precautions:

Use with caution in animals receiving digoxin, potassium-sparing diuretics, ACE inhibitors, and NSAIDs.

Lot No.:

Exp. Date:

Manufactured by Neogen Corporation

Lexington, KY 40511 USA • 859/254-1221

L592-91908

PRINCIPAL DISPLAY PANEL - 4 oz Bottle

NDC: 59051-9075-1

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NEOGEN.Vet

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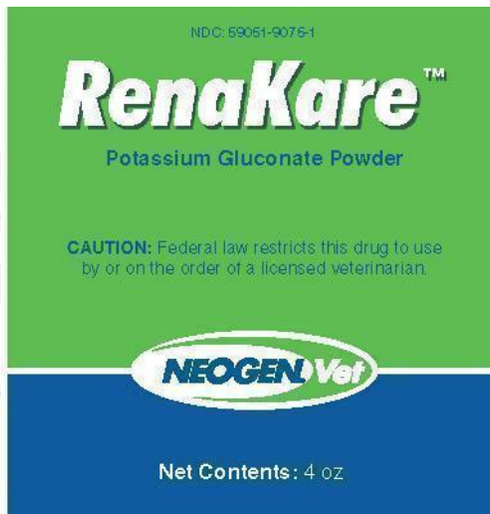
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 UPC



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RENAKARE			
potassium gluconate powder			
Product Information			
Product Type	PREScription ANIMAL DRUG	Item Code (Source)	NDC:59051-9075
Route of Administration	ORAL		
Active Ingredient/Active Moiety			
	Ingredient Name	Basis of Strength	Strength
	POTASSIUM GLUCONATE (UNII: 12H3K5QKN9) (POTASSIUM CATION - UNII:295O53K152)	POTASSIUM GLUCONATE	468 mg in 0.65 g
Product Characteristics			
Color	brown	Score	
Shape		Size	

Flavor		Imprint Code		
Contains				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:59051-9075-1	113 g in 1 BOTTLE, PLASTIC		
Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
unapproved drug other		10/24/2012		

Labeler - Neogen Corporation-Mercer Rd (042125879)

Establishment			
Name	Address	ID/FEI	Business Operations
Neogen Corporation-Mercer Rd		042125879	analysis, manufacture, label

Revised: 5/2013

Neogen Corporation-Mercer Rd