# **POTASSIUM CHLORIDE-** potassium chloride solution Genus Lifesciences Inc.

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#### HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use POTASSIUM CHLORIDE safely and effectively. See full prescribing information for POTASSIUM CHLORIDE.

#### POTASSIUM CHLORIDE oral solution

#### Initial U.S. Approval: 1948

INDICATIONS AND USAGE
Potassium Chloride is indicated for the treatment and prophylaxis of hypokalemia with or without metabolic alkalosis, in patients for whom dietary management with potassium-rich foods or diuretic dose reduction are insufficient. (1)
DOSAGE AND ADMINISTRATION
<ul> <li>Dilute prior to administration. (2.1, 5.1)</li> <li>Monitor serum potassium and adjust dosage accordingly (2.2, 2.3)</li> <li><i>Treatment of hypokalemia:</i></li> <li>Adults: Initial doses range from 40-100 mEq/day in 2-5 divided doses: limit doses to 40 mEq per dose. Total daily dose</li> </ul>
<ul> <li>Products: Initial doses range from 40-roo inEq/day in 2-5 divided doses. Initial doses to 40 inEq per dose. For a day dose should not exceed 200 mEq (2.2)</li> <li>Pediatric patients aged birth to 16 years old: 2-4 mEq/kg/day in divided doses; not to exceed 1 mEq/kg as a single dose</li> </ul>
• Pediatric patients aged birth to 10 years old. 2-4 mEq/kg/day in divided doses, not to exceed 1 mEq/kg as a single dose or 40 mEq whichever is lower; if deficits are severe or ongoing losses are great, consider intravenous therapy. Total daily dose should not exceed 100 mEq (2.3)
Maintenance or Prophylaxis of hypokalemia:
• Adults: Typical dose is 20 mEq per day (2.2)
• Pediatric patients aged birth to 16 years old: typical dose is 1 mEq/kg/day. Do not exceed 3 mEq/kg/day (2.3)
DOSAGE FORMS AND STRENGTHS
• Oral Solution: 10%; 1.3 mEq potassium per mL (3)
• Oral Solution: 20%; 2.6 mEq potassium per mL (3)
CONTRAINDICATIONS
Concomitant use with potassium sparing diuretics. (4)
WARNINGS AND PRECAUTIONS
• <u>Gastrointestinal Irritation</u> : Dilute before use, take with meals (5.1)
ADVERSE REACTIONS
Most common adverse reactions are nausea, vomiting, flatulence, abdominal pain/discomfort, and diarrhea. (6)
To report SUSPECTED ADVERSE REACTIONS, contact Pharm-Olam at 1-866-511-6754 or FDA at 1-800-FDA- 1088 or www.fda.gov/medwatch.
DRUG INTERACTIONS
<ul> <li>Potassium sparing diuretics: Avoid concomitant use (7.1)</li> </ul>
• Angiotensin converting enzyme inhibitors: Monitor for hyperkalemia (7.2)

Angiotensin receptor blockers: Monitor for hyperkalemia (7.3)

Revised: 5/2019

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#### FULL PRESCRIBING INFORMATION

#### **1 INDICATIONS AND USAGE**

Potassium Chloride is indicated for the treatment and prophylaxis of hypokalemia in patients for whom dietary management with potassium-rich foods or diuretic dose reduction are insufficient.

#### **2 DOSAGE AND ADMINISTRATION**

#### 2.1 Administration and Monitoring

#### Monitoring

Monitor serum potassium and adjust dosages accordingly. For treatment of hypokalemia, monitor potassium levels daily or more often depending on the severity of hypokalemia until they return to normal. Monitor potassium levels monthly to biannually for maintenance or prophylaxis.

The treatment of potassium depletion, particularly in the presence of cardiac disease, renal disease, or acidosis requires careful attention to acid-base balance, volume status, electrolytes, including magnesium, sodium, chloride, phosphate, and calcium, electrocardiograms and the clinical status of the patient. Correct volume status, acid-base balance and electrolyte deficits as appropriate.

#### Administration

Dilute the potassium chloride solution with at least 4 ounces of cold water [see Warnings and Precautions (5.1)].

Take with meals or immediately after eating.

If serum potassium concentration is  $<\!\!2.5$  mEq/L, use intravenous potassium instead of oral supplementation.

# 2.2 Adult Dosing

# Treatment of hypokalemia

Daily dose range from 40 to 100 mEq. Give in 2 to 5 divided doses: limit doses to 40 mEq per dose. The total daily dose should not exceed 200 mEq in a 24 hour period.

# Maintenance or Prophylaxis

Typical dose is 20 mEq per day. Individualize dose based upon serum potassium levels.

Studies support the use of potassium replacement in digitalis toxicity. When alkalosis is present, normokalemia and hyperkalemia may obscure a total potassium deficit. The advisability of use of potassium replacement in the setting of hyperkalemia is uncertain.

# 2.3 Pediatric Dosing

# Treatment of hypokalemia

Pediatric patients aged birth to 16 years old: The initial dose is 2 to 4 mEq/kg/day in divided doses; do not exceed as a single dose 1 mEq/kg or 40 mEq, whichever is lower; maximum daily doses should not exceed 100 mEq. If deficits are severe or ongoing losses are great, consider intravenous therapy.

# Maintenance or Prophylaxis

Pediatric patients aged birth to 16 years old: Typical dose is 1 mEq/kg/day. Do not exceed 3 mEq/kg/day.

# **3 DOSAGE FORMS AND STRENGTHS**

Oral Solution 10%: 1.3 mEq potassium per mL.

Oral Solution 20%: 2.6 mEq potassium per mL.

# **4 CONTRAINDICATIONS**

Potassium chloride is contraindicated in patients on potassium sparing diuretics.

# **5 WARNINGS AND PRECAUTIONS**

## 5.1 Gas trointes tinal Irritation

May cause gastrointestinal irritation if administered undiluted. Increased dilution of the solution and taking with meals may reduce gastrointestinal irritation [*see Dosage and Administration (2.1)*].

# **6 ADVERSE REACTIONS**

The most common adverse reactions to oral potassium salts are nausea, vomiting, flatulence, abdominal pain/discomfort, and diarrhea.

# 7 DRUG INTERACTIONS

# 7.1 Potassium-Sparing Diuretics

Use with potassium-sparing diuretics can produce severe hyperkalemia. Avoid concomitant use.

### 7.2 Angiotensin-Converting Enzyme Inhibitors

Use with angiotensin converting enzyme (ACE) inhibitors produces potassium retention by inhibiting aldosterone production. Potassium supplements should be given to patients receiving ACE inhibitors only with close monitoring.

#### 7.3 Angiotensin Receptor Blockers

Use with angiotensin receptor blockers (ARBs) produces potassium retention by inhibiting aldosterone production. Potassium supplements should be given to patients receiving ARBs only with close monitoring.

## **8 USE IN SPECIFIC POPULATIONS**

#### 8.1 Pregnancy

Pregnancy Category C

Animal reproduction studies have not been conducted with potassium chloride. It is unlikely that potassium supplementation that does not lead to hyperkalemia would have an adverse effect on the fetus or would affect reproductive capacity.

#### 8.2 Nursing Mothers

The normal potassium ion content of human milk is about 13 mEq per liter. Since oral potassium becomes part of the body potassium pool, so long as body potassium is not excessive, the contribution of potassium chloride supplementation should have little or no effect on the level in human milk.

#### 8.3 Pediatric Use

The safety and effectiveness of potassium chloride have been demonstrated in children with diarrhea and malnutrition from birth to 16 years.

#### 8.4 Geriatric Use

Clinical studies of Potassium Chloride did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects. Other reported clinical experience has not identified differences in responses between the elderly and younger patients. In general, dose selection for an elderly patient should be cautious, usually starting at the low end of the dosing range, reflecting the greater frequency of decreased hepatic, renal, or cardiac function, and of concomitant disease or other drug therapy.

This drug is known to be substantially excreted by the kidney, and the risk of toxic reactions to this drug may be greater in patients with impaired renal function. Because elderly patients are more likely to have decreased renal function, care should be taken in dose selection, and it may be useful to monitor renal function.

#### **10 OVERDOSAGE**

#### 10.1 Symptoms

The administration of oral potassium salts to persons with normal excretory mechanisms for potassium rarely causes serious hyperkalemia. However, if excretory mechanisms are impaired or if potassium is administered too rapidly potentially fatal hyperkalemia can result.

Hyperkalemia is usually asymptomatic and may be manifested only by an increased serum potassium concentration (6.5–8.0 mEq/L) and characteristic electrocardiographic changes (peaking of T-waves,

loss of P-waves, depression of S-T segment, and prolongation of the QT-interval). Late manifestations include muscle paralysis and cardiovascular collapse from cardiac arrest (9–12 mEq/L).

## 10.2 Treatment

Treatment measures for hyperkalemia include the following:

- 1. Monitor closely for arrhythmias and electrolyte changes.
- 2. Eliminate foods and medications containing potassium and of any agents with potassium-sparing properties such as potassium-sparing diuretics, ARBS, ACE inhibitors, NSAIDS, certain nutritional supplements and many others.
- 3. Administer intravenous calcium gluconate if the patient is at no risk or low risk of developing digitalis toxicity.
- 4. Administer intravenously 300 to 500 mL/hr of 10% dextrose solution containing 10 to 20 units of crystalline insulin per 1000 mL.
- 5. Correct acidosis, if present, with intravenous sodium bicarbonate.
- 6. Use exchange resins, hemodialysis, or peritoneal dialysis.

In patients who have been stabilized on digitalis, too rapid a lowering of the serum potassium concentration can produce digitalis toxicity.

# **11 DESCRIPTION**

Potassium Chloride is a white crystalline or colorless solid. It is soluble in water and slightly soluble in alcohol. Chemically, Potassium Chloride is K-Cl with a molecular mass of 74.55.

Oral Solution: 10%: Each 15 mL of solution contains 1.5 g of potassium chloride, USP and the following inactive ingredients: citric acid anhydrous, FD&C Yellow #6, glycerin, natural/artificial orange flavor, purified water, sodium benzoate, sodium citrate dihydrate, sucralose.

Oral Solution 20%: Each 15 mL of solution contains 3.0 g of potassium chloride, USP and the following inactive ingredients: citric acid anhydrous, FD&C Yellow #6, glycerin, natural/artificial orange flavor, purified water, sodium benzoate, sodium citrate dihydrate, sucralose.

# **12 CLINICAL PHARMACOLOGY**

# 12.1 Mechanism of Action

The potassium ion (K+) is the principal intracellular cation of most body tissues. Potassium ions participate in a number of essential physiological processes including the maintenance of intracellular tonicity; the transmission of nerve impulses; the contraction of cardiac, skeletal, and smooth muscle; and the maintenance of normal renal function.

The intracellular concentration of potassium is approximately 150 to 160 mEq per liter. The normal adult plasma concentration is 3.5 to 5 mEq per liter. An active ion transport system maintains this gradient across the plasma membrane.

Potassium is a normal dietary constituent, and under steady-state conditions the amount of potassium absorbed from the gastrointestinal tract is equal to the amount excreted in the urine. The usual dietary intake of potassium is 50 to 100 mEq per day.

# 12.3 Pharmacokinetics

Based on published literature, the rate of absorption and urinary excretion of potassium from KCl oral solution were higher during the first few hours after dosing relative to modified release KCl products. The bioavailability of potassium, as measured by the cumulative urinary excretion of K+ over a 24 hour post dose period, is similar for KCl solution and modified release products.

### **16 HOW SUPPLIED/STORAGE AND HANDLING**

Potassium Chloride Oral Solution, is an orange solution available in two strengths as follows:

10%: 20 mEq/15 mL oral solution

NDC# 64950-320-47 Bottle of 473 mL NDC# 64950-320-99 Bulk Drum of 30 gallons

20%: 40 mEq/15 mL oral solution

NDC# 64950-322-47 Bottle of 473 mL

### Storage

Store at Controlled Room Temperature, 25°C (77°F); excursions are permitted to 15° - 30°C (59° - 86°F).

Dispense in a tight, light-resistant container as defined in the USP.

PROTECT from LIGHT and FREEZING.

Rx only

Manufactured by: Genus Lifesciences Inc. Allentown, PA 18102

Revision 5/2019 r1

## PRINCIPAL DISPLAY PANEL - 30 gallon Drum Label

Each 15 mL (tablespoon) contains: Potassium Chloride, USP 20 mEq

Inactive ingredients: citric acid, FD&C Yellow #6, glycerin, natural/artificial orange flavor, purified water, sodium benzoate, sodium citrate dihydrate, sucralose.

Dosage and Administration: See accompanying prescribing information.

Store at 25°C (77°F); excursions permitted to 15°-30°C (59°-86°F).

Protect from Light and Freezing

KEEP THIS AND ALL MEDICA-TION OUT OF THE REACH OF CHILDREN.

Manufactured by: Genus Lifesciences Inc. Allentown, PA 18102

#### Rev. 11/18

NDC 64950-320-99

Potassium Chloride Oral Solution, USP, 10%

20 mEq per 15 mL

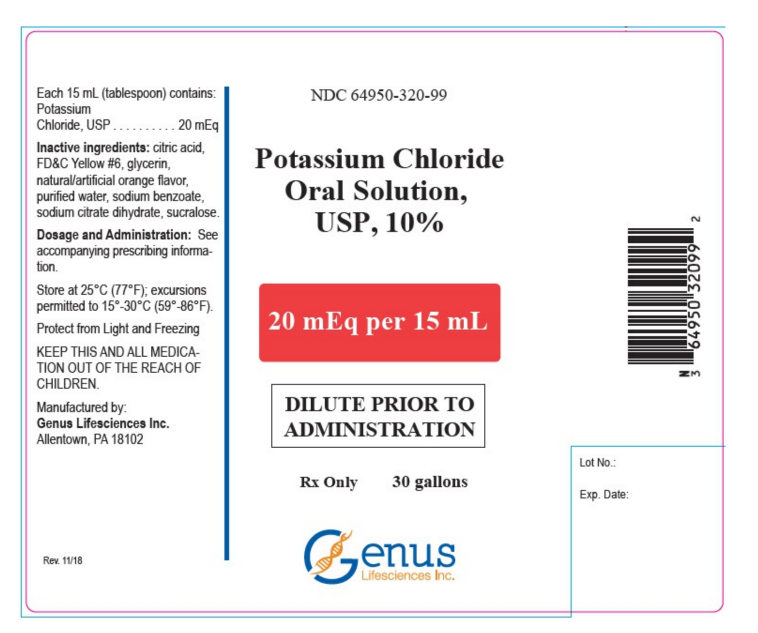
DILUTE PRIOR TO ADMINISTRATION

Rx Only 30 gallons

Genus Lifesciences Inc.

Lot No.:

Exp. Date:



#### PRINCIPAL DISPLAY PANEL - 473 mL Bottle Label

NDC 64950-322-47

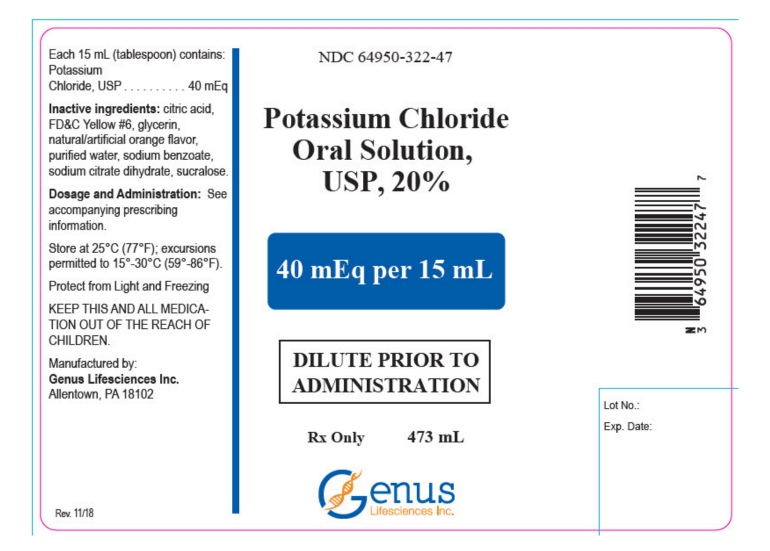
Potassium Chloride Oral Solution, USP, 20%

40 mEq per 15 mL

DILUTE PRIOR TO ADMINISTRATION

Rx Only 473 mL

Genus Lifesciences Inc.



# POTASSIUM CHLORIDE potassium chloride solution

**Product Information** 

Product Type

HUMAN PRESCRIPTION DRUG

Item Code (Source)

NDC:64950-320

ORAL
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Active Ingredier	nt/Active Moiety				
	Ingredient Name Basis of Strength				th
Potassium Chloride	(UNII: 660 YQ98110) (POTASSIUM CATION - UNII:295053)	K152)	Potassium Chloride	20 meq in	15 mL
Inactive Ingredi	ents				
	Ingredient Name			Streng	th
ANHYDRO US CITRI	C ACID (UNII: XF417D3PSL)				
FD&C YELLOW NO	<b>6</b> (UNII: H77VEI93A8)				
Glycerin (UNII: PDC6	A3C0OX)				
Sodium Benzoate (U	NII: OJ245FE5EU)				
SODIUM CITRATE, U	J <b>NSPECIFIED FORM</b> (UNII: 1Q73Q2JULR)				
Sucralose (UNII: 96K	6UQ3ZD4)				
Water (UNII: 059QF0	KO0R)				
<b>Product Charact</b>	eristics				
Color	YELLOW		Score		
Shape	Size				
Flavor	ORANGE (Natural & Artificial)	Imprint Code			
Contains					
Packaging					
# Item Code	Package Description	N	larketing Start Date	Marketing Date	End
1 NDC:64950-320- 47	473 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	n 02/1	5/2015	09/07/2015	
2 NDC:64950-320- 99	113562 mL in 1 DRUM; Type 0: Not a Combination Product	0 1/2	0 1/2 1/20 16		
Marketing Information					
Marketing Categor		Manko	ing Start Data	Iarkating End	Data
			-	larketing End	Date
NDA	NDA206814	02/15/201			

# **POTASSIUM CHLORIDE**

potassium chloride solution

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:64950-322
Route of Administration	ORAL		

Active Ingreui	ent/Active Moie	L <b>y</b>			
Ingredient Name Basis of Strength					gth Strength
Potassium Chloride (UNII: 660 YQ98 I10) (POTASSIUM CATION - UNII:295053K152) Potassium Chloride					e 40 meq in 15 m
Inactive Ingree	lients				
		Ingredient Name			Strength
ANHYDRO US CITI	IC ACID (UNII: XF41	7D3PSL)			
FD&C YELLOW N	<b>D.6</b> (UNII: H77VEI93	A8)			
<b>Glycerin</b> (UNII: PDO	6A3C0OX)				
	UNII: OJ245FE5EU)				
SO DIUM CITRATE	UNSPECIFIED FOF	<b>RM</b> (UNII: 1Q73Q2JULR)			
	ucralose (UNII: 96K6UQ3ZD4)				
Water (UNII: 059Q)	0KO0R)				
Product Chara				_	
Color	YELLOW	YELLOW Score			
Shape		Size			
Flavor	ORANGE (Natura	ll & Artificial)		Imprint Code	
Contains					
Packaging					
# Item Code	]	Package Description		Marketing Start Date	Marketing End Date
1 NDC:64950-322- 47	473 mL in 1 BOTTL Product	E, PLASTIC; Type 0: Not a Combination	on C	2/15/2015	09/07/2015
Marketing I	nformation				
Marketing I Marketing Categ		Number or Monograph Citation	Mar	keting Start Date	Marketing End Dat

Labeler - Genus Lifesciences Inc. (113290444)

Establishment			
Name	Address	ID/FEI	Business Operations
Genus Lifesciences Inc.		113290444	MANUFACTURE(64950-320, 64950-322), PACK(64950-320), LABEL(64950-320)

Revised: 7/2019

Genus Lifesciences Inc.