

SELECT BRAND ANTI-NAUSEA- phosphorated carbohydrate solution

L&R Distributors

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](#).

Select Brand Anti-Nausea Liquid

DRUG FACTS

Active ingredients

Phosphorated carbohydrate solution*

*Each 5 mL contains:

- 3.74g Total sugar
- 21.5mg Phosphoric acid

Purpose

Upset stomach reliever

Uses

For relief of nausea due to upset stomach from intestinal flu, stomach flu, and food or drink indiscretions.

Warnings

- this product contains fructose and should not be taken by persons with hereditary fructose intolerance (HFI).

As with any drug, if you are pregnant or nursing a baby, seek the advice of a health professional before using this product.

Stop use and ask a doctor if

- symptoms persist, return or get worse

KEEP THIS AND ALL MEDICATIONS OUT OF THE REACH OF CHILDREN.

In case of accidental overdose, seek professional assistance or contact a Poison Control Center right away.

DOSAGE: Administer dosage below every 15 minutes or until distress subsides.

Adults - One or two tablespoons.

Children 2 to 12 - One or two teaspoons.

Other information

Store at room temperature

Inactive ingredients

FD&C Red No 40, flavor, glycerin, methylparaben, and purified water.

PACKAGE LABEL

Lable

NDC 15127-291-26

select brand
the lower price name brand

Anti-Nausea Liquid

for Nausea Associated with upset Stomach

Cherry Flavor

Compare to Emetrol®*

4 FL OZ (118 mL)

DO NOT USE IF PRINTED SEAL UNDER CAP IS TORN OR MISSING

INDICATIONS: For relief of nausea due to upset stomach from intestinal flu, stomach flu, and food or drink indiscretions.

DOSAGE: Administer dosage below every 15 minutes or until distress subsides.
Adults- One or two teaspoons.
Children 2 to 12- One or two teaspoons.

CAUTION: Not to be taken for more than one hour (5 doses) without consulting a physician. If nausea continues or recurs frequently, consult a physician.

FOR MAXIMUM EFFECTIVENESS NEVER DILUTE ANTI-NAUSEA or drink fluids of any kind immediately before or after taking.

WARNINGS: this product contains fructose and should not be taken by persons with hereditary fructose intolerance (HFI).
As with any drug, if you are pregnant or nursing a baby, seek the advice of a health professional before using this product.

THIS PRODUCT CONTAINS SUGAR AND SHOULD NOT BE TAKEN BY DIABETICS EXCEPT UNDER THE ADVICE AND SUPERVISION OF A PHYSICIAN

In case of accidental overdose, seek professional assistance or contact a poison control center immediately.

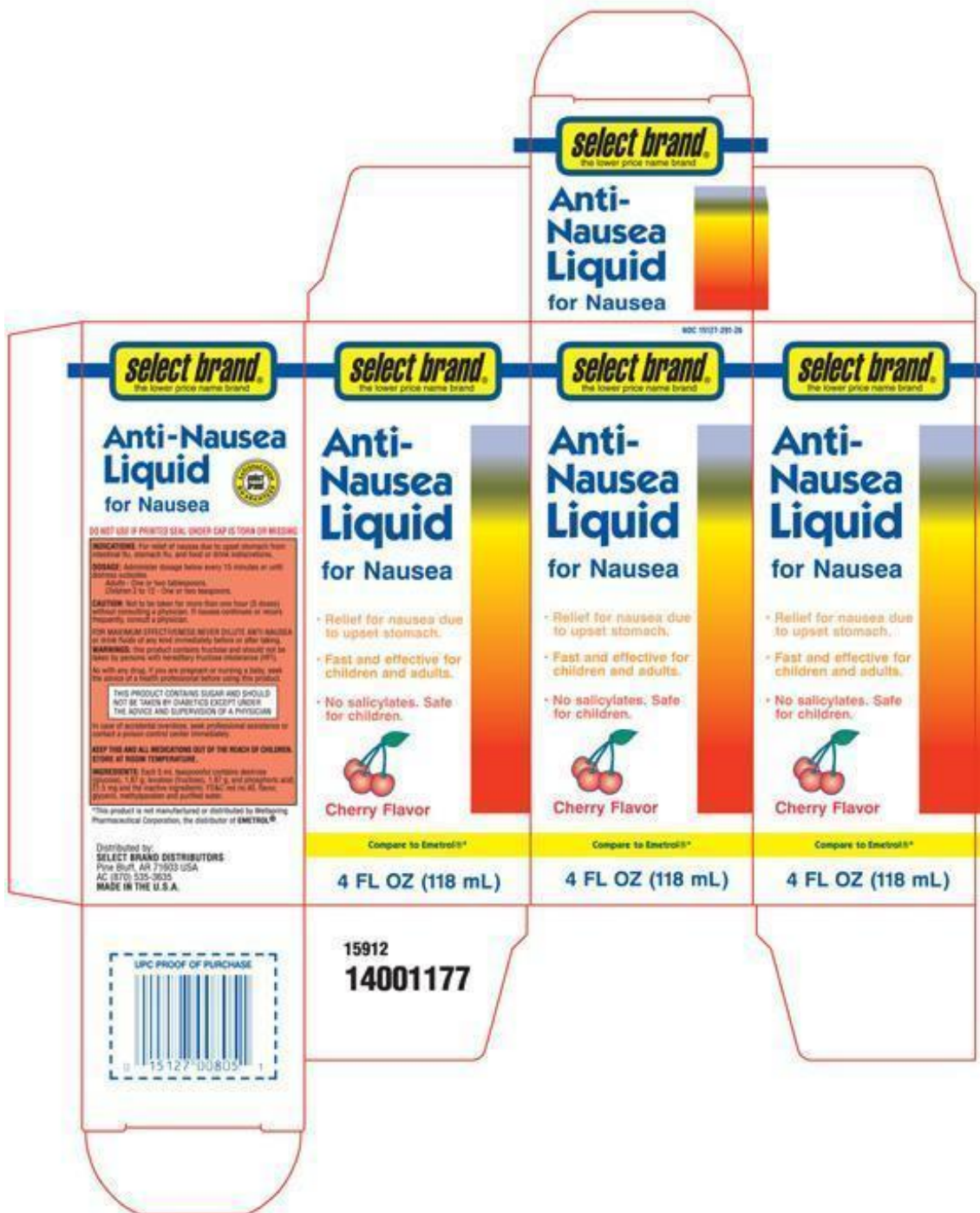
KEEP THIS AND ALL MEDICATIONS OUT OF THE REACH OF CHILDREN. STORE AT ROOM TEMPERATURE.

INGREDIENTS: Each 5 mL teaspoonful contains dextrose (glucose), 1.87 g; levulose (fructose), 1.87 g; and phosphoric acid, 21.5 mg and the inactive ingredients: FD&C red no.40, flavor, glycerin, methylparaben and purified water.

*This product is not manufactured or distributed by **14001178** Wellspring Pharmaceutical Corporation, the distributor of EMETROL®.
Distributed by: **SELECT BRAND DISTRIBUTORS**
Pine Bluff, AR 71603 U.S.A.
AC (870) 535-3635
MADE IN THE USA

0 15127 00805 1

Carton



SELECT BRAND ANTI-NAUSEA

phosphorated carbohydrate solution

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:15127-291
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SUCROSE (UNII: C151H8M554) (SUCROSE - UNII:C151H8M554)	SUCROSE	3.74 g in 5 mL
PHOSPHORIC ACID (UNII: E4GA8884NN) (PHOSPHORIC ACID - UNII:E4GA8884NN)	PHOSPHORIC ACID	21.5 mg in 5 mL

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
GLYCERIN (UNII: PDC6A3C0OX)	
METHYLPARABEN (UNII: A2I8C7H9T)	
FD&C RED NO. 40 (UNII: WZB9127XOA)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:15127-291-26	118 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	09/29/2014	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		01/01/2008	

Labeler - L&R Distributors (012578514)

Registrant - Humco Holding Group, Inc. (825672884)

Establishment

Name	Address	ID/FEI	Business Operations
Humco Holding Group, Inc.		825672884	label(15127-291) , manufacture(15127-291)

Revised: 12/2020

L&R Distributors